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=> s redispersible

L1 687 REDISPERSIBLE

=> s redispersion

1070 REDISPERSION

8 REDISPERSIONS

L2 1077 REDISPERSION

(REDISPERSION OR REDISPERSIONS)

=> s ZnO

103378 ZNO

11 ZNOS

L3 103382 ZNO

(ZNO OR ZNOS)

=> s zinc oxide

638647 ZINC

139 ZINCS

638672 ZINC

(ZINC OR ZINCS)

1796554 OXIDE

349434 OXIDES

1895107 OXIDE

(OXIDE OR OXIDES)

L4 103202 ZINC OXIDE

(ZINC(W)OXIDE)

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L5 26 (L1 OR L2) AND (L3 OR L4)

=> display all L5 1-26 ibib abs

L5 ANSWER 1 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

AN 2007:970296 CAPLUS

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Entered STN: 31 Aug 2007
ΕD
     Low-viscosity water-in-oil cosmetic emulsions containing water-soluble
ΤI
     UV-absorbing sulfonic acids, titania or zinc oxide,
     and silicone oils
ΙN
     Yamaguchi, Kazuhiro
PA
     Shiseido Co., Ltd., Japan
     Jpn. Kokai Tokkyo Koho, 15pp.
SO
     CODEN: JKXXAF
DT
     Patent
LA
     Japanese
CC
     62-4 (Essential Oils and Cosmetics)
FAN.CNT 1
     PATENT NO.
                                DATE APPLICATION NO.
                        KIND
                                                                  DATE
                        ____
                                            -----
    JP 2007217393
                         Α
                                20070830 JP 2006-43159
                                                                   20060220
PRAI JP 2006-43159
                                20060220
CLASS
PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES
A61K0008-49 [I,A]; A61K0008-30 [I,C*]; A61K0008-20
                        [I,A]; A61K0008-23 [I,A]; A61K0008-06 [I,A];
                        A61K0008-04 [I,C*]
                       4C083/AA122; 4C083/AB051; 4C083/AB052; 4C083/AB082;
                 FTERM
                        4C083/AB211; 4C083/AB212; 4C083/AB241; 4C083/AB242;
                        4C083/AB331; 4C083/AB332; 4C083/AB351; 4C083/AC022;
                        4C083/AC102; 4C083/AC122; 4C083/AC342; 4C083/AC372;
                        4C083/AC422; 4C083/AC542; 4C083/AC642; 4C083/AC662; 4C083/AC851; 4C083/AC852; 4C083/AD151; 4C083/AD152;
                        4C083/AD162; 4C083/AD172; 4C083/AD662; 4C083/BB46;
                        4C083/CC19; 4C083/DD32; 4C083/EE01; 4C083/EE06;
                        4C083/EE17
AΒ
    Title cosmetics, which show good storage stability, no stickiness, and
    smoothly-spreading property and are stirred with a steel ball just before
    the use, contain (a) water-soluble UV absorbers having sulfo group, (b) TiO2
    and/or ZnO, (c) \geq10% silicone oils, (d) H2O, and optional
     (e) \geq 1 selected from NaCl, MgCl2, Na2SO4, and MgSO4 to improve
    emulsification stability. Thus, a W/O sunscreen emulsion was formulated
    containing decamethylcyclopentasiloxane, dimethicone, dimethicone-treated
    ZnO, H2O, Na phenylbenzimidazole-5-sulfonate, and NaCl. The
    emulsion packed in a container with a stainless steel ball was shaken 10
    times to show uniformly dispersed state. The emulsion was stored at
    50° for 1 mo to show no changes in the appearance, the odor, and
    the touch in application to skin.
    sunscreen emulsion UV absorbing sulfonic acid titania silicone oil; phenylbenzimidazolesulfonate UV absorber zinc oxide
ST
    silicone sunscreen emulsion
ΙT
    Cosmetic emulsions
    Sunscreens
        (easily-redispersible low-viscosity W/O cosmetic emulsions
       containing water-soluble UV-absorbing sulfonic acids, TiO2 or ZnO,
       silicone oils, and H2O)
ΙT
    Cvclosiloxanes
    Polysiloxanes
    RL: COS (Cosmetic use); BIOL (Biological study); USES (Uses)
        (easily-redispersible low-viscosity W/O cosmetic emulsions
       containing water-soluble UV-absorbing sulfonic acids, TiO2 or ZnO,
       silicone oils, and H2O)
    541-02-6, Decamethylcyclopentasiloxane 1314-13-2, Zinc
IT
            5997-53-5 7732-18-5, Water 9006-65-9, Dimethicone
    13463-67-7, Titania 27503-81-7 170864-82-1 180898-37-7, Neo Heliopan
    RL: COS (Cosmetic use); BIOL (Biological study); USES (Uses)
        (easily-redispersible low-viscosity W/O cosmetic emulsions
```

containing water-soluble UV-absorbing sulfonic acids, TiO2 or ZnO, silicone oils, and H2O) 7487-88-9, Magnesium sulfate 7647-14-5, Sodium chloride ΤТ 7757-82-6, Sodium sulfate 7786-30-3, Magnesium chloride RL: COS (Cosmetic use); BIOL (Biological study); USES (Uses) (stabilizer; easily-redispersible low-viscosity W/O cosmetic emulsions containing water-soluble UV-absorbing sulfonic acids, TiO2 or ZnO, silicone oils, and H2O) ACCESSION NUMBER: 2007:970296 CAPLUS TITLE: Low-viscosity water-in-oil cosmetic emulsions containing water-soluble UV-absorbing sulfonic acids, titania or zinc oxide, and silicone oils INVENTOR(S): Yamaguchi, Kazuhiro PATENT ASSIGNEE(S): Shiseido Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 15pp. CODEN: JKXXAF DOCUMENT TYPE: Patent LANGUAGE: Japanese FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. -----____ ----------JP 2007217393 A 20070830 JP 2006-43159 20060220 JP 2006-43159 20060220 PRIORITY APPLN. INFO.: Title cosmetics, which show good storage stability, no stickiness, and smoothly-spreading property and are stirred with a steel ball just before the use, contain (a) water-soluble UV absorbers having sulfo group, (b) TiO2 and/or ZnO, (c) ≥10% silicone oils, (d) H2O, and optional (e) ≥ 1 selected from NaCl, MgCl2, Na2SO4, and MgSO4 to improve emulsification stability. Thus, a W/O sunscreen emulsion was formulated containing decamethylcyclopentasiloxane, dimethicone, dimethicone-treated ZnO, H2O, Na phenylbenzimidazole-5-sulfonate, and NaCl. The emulsion packed in a container with a stainless steel ball was shaken 10 times to show uniformly dispersed state. The emulsion was stored at 50° for 1 mo to show no changes in the appearance, the odor, and the touch in application to skin. ANSWER 2 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN 1.5 2007:816790 CAPLUS AN 147:199722 DN Entered STN: 27 Jul 2007 ED ΤI Process for the synthesis of nanocrystalline phosphor particles and nanocrystalline phosphor particles obtainable by this process ΙN Feldmann, Claus; Buehler, Gunnar PA Universitaet Karlsruhe, Germany SO PCT Int. Appl., 34pp. CODEN: PIXXD2 DT Patent LA German 73-5 (Optical, Electron, and Mass Spectroscopy and Other Related Properties) Section cross-reference(s): 9, 42, 78 FAN.CNT 1 PATENT NO. APPLICATION NO. KIND DATE ----------____ _____ WO 2007082663 A1 20070726 WO 2007-EP175 20070110 PΙ W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,

GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT,

TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW

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             IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ,
             CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH,
             GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU, TJ, TM
                       A1
     DE 102006001414
                                20070816
                                          DE 2006-102006001414 20060111
PRAI DE 2006-102006001414 A
                                20060111
CLASS
 PATENT NO.
                 CLASS PATENT FAMILY CLASSIFICATION CODES
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 WO 2007082663 IPCI
                        C09K0011-08 [I,A]; C09K0011-02 [I,A]
 DE 102006001414 IPCI
                        C09K0011-00 [I,A]; C09K0011-08 [I,A]; C08K0003-00
                        [I,A]; A61K0049-00 [I,A]; D21H0021-30 [I,A];
                        D21H0021-14 [I,C*]; D21H0021-40 [I,A]; B05D0005-06
                        [I,A]; C03C0004-12 [I,A]; C03C0004-00 [I,C*];
                        C03C0017-22 [I,A]; C04B0035-00 [I,A]; C04B0041-85 [I,A]
                 ECLA
                        C09K011/02; C09K011/08
AB
     Methods for the targeted synthesis of inorg. phosphor particles which have
     a particle diameter of 1-100 nm, a virtually monodisperse size distribution
     in the range of \pm 20\% and, depending on the class of material selected,
     a quantum yield of ≥20% which entail reacting ≥1 metal
     precursor with \geq 1 precipitating reagent at -50^{\circ} to 200° in a
     first solvent or solvent mixture; thermally treating the resulting mixture at
     150-700° for 5 s to 30 min in a second solvent or solvent mixture
     that includes ≥1 ionic liquid; and isolating and/or purifying the
     phosphor particles are described in which the metal precursor is chosen
     from halides, nitrates, sulfates, hydrogen sulfates, acetates,
     acetylacetonates, oxalates, carbonates, hydrogen carbonates, phosphates,
     hydrogen phosphates, dihydrogen phosphates, hydrides, alcoholates, and/or
     metalorg. compds., and the precipitating reagent is chosen from hydroxides,
     phosphates, carbonates, oxalates, sulfates, sulfides, halides, ammonia,
     hydroxylamines, amines, amides, imides, and/or water. Inorg. phosphor
     particles obtainable by this process and also their use for coating or
     embedding in a substrate selected from the group consisting of a paper,
     polymer, glass, metal or ceramic substrate, for example for diagnostics,
     therapy, as contrast agent, as marking, warning feature, for illumination,
     and for data storage and/or data reproduction. A particular advantage of the
     phosphor particles which can be obtained according to the invention is
     that the particles have a surface finish which allows agglomerate-free
     redispersion in polar and nonpolar solvents or dispersion media.
     The choice of the dispersion medium allows the physicochem. properties of
     the phosphor particles which can be obtained according to the invention to
     be specifically matched to the requirements of a particular application.
ST
     nanocryst phosphor particle prepn; diagnostics nanocryst phosphor
     particle; therapy nanocryst phosphor particle; contrast agent nanocryst
     phosphor particle; marking nanocryst phosphor particle; illumination
     nanocryst phosphor particle; data storage nanocryst phosphor particle
IT
     Imaging agents
        (contrast; synthesis of nanocryst. phosphor particles and the
        nanocryst. phosphor particles and their uses)
IT
     Drugs
     Marking
     Nanocrystals
     Phosphors
     Recording materials
        (synthesis of nanocryst. phosphor particles and the nanocryst. phosphor
       particles and their uses)
TΤ
    Amides, processes
     Amines, processes
     Carbonates, processes
     Coordination compounds
     Halides
     Imides
    Metal alkoxides
    Nitrates, processes
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Phosphates, processes
     Sulfates, processes
    Sulfides, processes
    RL: PEP (Physical, engineering or chemical process); RCT (Reactant); PROC
     (Process); RACT (Reactant or reagent)
        (synthesis of nanocryst. phosphor particles and the nanocryst. phosphor
        particles and their uses)
ΙT
    7429-90-5P, Aluminum, uses
                                 7429-91-6P, Dysprosium, uses
                                                                7439-92-1P,
    Lead, uses
                 7439-95-4P, Magnesium, uses
                                               7439-96-5P, Manganese, uses
    7440-00-8P, Neodymium, uses 7440-02-0P, Nickel, uses
                                                            7440-03-1P,
                    7440-10-0P, Praseodymium, uses 7440-19-9P, Samarium,
    Niobium, uses
           7440-21-3P, Silicon, uses 7440-22-4P, Silver, uses
                                                                 7440-23-5P,
                   7440-27-9P, Terbium, uses
    Sodium, uses
                                               7440-30-4P, Thulium, uses
                            7440-32-6P, Titanium, uses 7440-36-0P, Antimony,
    7440-31-5P, Tin, uses
           7440-45-1P, Cerium, uses
                                      7440-50-8P, Copper, uses
                                                                 7440-52-0P,
                   7440-53-1P, Europium, uses
    Erbium, uses
                                               7440-55-3P, Gallium, uses
    7440-64-4P, Ytterbium, uses
                                  7440-66-6P, Zinc, uses
                                                           7440-69-9P,
                    13708-63-9P, Terbium fluoride 13765-25-8P, Europium
    Bismuth, uses
               113671-38-8P, Silicon oxide (SiO0-2)
    RL: IMF (Industrial manufacture); MOA (Modifier or additive use); TEM
     (Technical or engineered material use); PREP (Preparation); USES (Uses)
        (phosphors doped with; synthesis of nanocryst. phosphor particles and
        the nanocryst. phosphor particles and their uses)
ΙT
    1305-78-8P, Calcium oxide, uses
                                     1314-13-2P, Zinc oxide
             1314-36-9P, Yttria, uses 1314-96-1P, Strontium sulfide
    1314-98-3P, Zinc sulfide, uses
                                    1344-28-1P, Alumina, uses
                                                                 7631-86-9P,
    Silica, uses
                   7758-87-4P, Calcium phosphate (Ca3(PO4)2)
                                                               7778-18-9P,
    Calcium sulfate (CaSO4)
                              7779-90-0P, Zinc phosphate (Zn3(PO4)2)
    7783-40-6P, Magnesium difluoride
                                       7789-17-5P, Cesium iodide 7789-24-4P,
    Lithium fluoride, uses 7789-75-5P, Calcium fluoride, uses
                                                                  7790-75-2P,
    Calcium tungsten oxide (CaWO4)
                                     10101-39-0P
                                                   10377-51-2P, Lithium iodide
    12003-86-0P, Yttrium aluminate (YAlO3)
                                             12004-04-5P, Barium aluminate
                12005-21-9P, YAG 12007-60-2P, Lithium borate (Li2B407)
     (BaAl204)
    12024-21-4P, Gallium oxide
                                 12024-36-1P, Gadolinium gallium oxide
                  12027-88-2P, Yttrium silicate (Y2SiO5)
    (Gd3Ga5012)
                                                           12031-43-5P,
    Lanthanum oxide sulfide (La2O2S) 12031-63-9P, Lithium niobate
    12032-36-9P, Magnesium sulfide
                                    12143-49-6P, Yttrium tantalate (YTaO4)
    12159-91-0P, Germanium magnesium fluoride oxide (Ge2Mg8F2O11)
    12233-56-6P, Bismuth germanium oxide (Bi4Ge3012)
                                                       12254-04-5P, Aluminum
    barium magnesium oxide (Al10BaMgO17)
                                           12339-07-0P, Gadolinium oxide
    sulfide (Gd2O2S)
                       12340-04-4P, Yttrium oxide sulfide (Y202S)
    12442-27-2P, Cadmium zinc sulfide ((Cd, Zn)S) 12592-70-0P, Strontium
    gallium sulfide (SrGa2S4)
                                13466-21-2P, Barium phosphate (Ba2P2O7)
    13566-12-6P, Yttrium vanadate
                                   13568-56-4P, Lutetium vanadate (LuVO4)
    13573-11-0P, Magnesium tungstate (MgWO4)
                                               13597-55-2P, Strontium silicate
    (Sr2SiO4)
                13597-65-4P, Zinc silicate (Zn2SiO4)
                                                      13628-52-9P, Gadolinium
                       13709-38-1P, Lanthanum trifluoride
    vanadate (GdVO4)
                                                           13709-49-4P,
    Yttrium trifluoride
                          13718-55-3P, Barium chloride fluoride (BaClF)
    13759-29-0P, Yttrium chloride oxide (YClO)
                                                13776-74-4P, Magnesium
    silicate (MgSiO3)
                       13778-49-9P, Barium silicate (Ba2SiO4)
                                                                13778-59-1P
    13812-81-2P, Strontium phosphate (Sr2P2O7)
                                                 13875-40-6P, Lanthanum oxide
    bromide (LaOBr)
                     13968-67-7P, Barium silicate (BaSi2O5) 14118-26-4P,
    Sodium lanthanum fluoride (NaLaF4)
                                         15640-94-5P, Sodium gadolinium
    fluoride (NaGdF4)
                        20548-54-3P, Calcium sulfide 21669-04-5P, Barium
    bromide fluoride (BaBrF)
                               23108-36-3P, Yttrium lithium fluoride (YLiF4)
    25617-97-4P, Gallium nitride
                                  26874-36-2P, Barium yttrium fluoride
              26916-94-9P, Lithium lutetium fluoride (LiLuF4)
    (BaYF5)
                                                                28042-61-7P,
    Potassium magnesium fluoride (KMgF3)
                                         33846-79-6P, Barium yttrium
    fluoride (BaY2F8)
                        35361-71-8P, Lithium strontium aluminum fluoride
    (LiSrAlF6)
                 35362-46-0P, Lithium calcium aluminum fluoride (LiCaAlF6)
    37276-56-5P, Calcium strontium chloride phosphate (CaSr9Cl2(PO4)6)
    55134-50-4P, Aluminum barium magnesium oxide (Al16BaMg2027)
                                                                  69142-81-0P
    75529-26-9P, Gadolinium magnesium borate (GdMgB5010)
                                                           75535-31-8P.
    Calcium chloride fluoride phosphate (Ca5(Cl,F)(PO4)3)
                                                            76125-60-5P,
                                      79304-71-5P, Lanthanum silicon nitride
    Strontium aluminate (Sr4Al14025)
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(LaSi3N5)
                82992-94-7P, Calcium strontium sulfide ((Ca,Sr)S)
     106804-21-1P, Magnesium strontium phosphate ((Mg,Sr)3(PO4)2)
     115968-61-1P, Vanadium yttrium oxide phosphate ([V,O,(PO4)]Y)
     119537-26-7P, Calcium magnesium sulfide ((Ca,Mg)S)
                                                          122656-71-7P, Barium
     bromide chloride fluoride (BaBr0.5Cl0.5F)
                                                 124676-67-1P, Gadolinium
     yttrium borate ((Gd,Y)(BO3))
                                    126344-47-6P, Magnesium zinc fluoride
     ((Mg, Zn)F2)
                   127734-77-4P
                                 144419-68-1P, Aluminum barium cerium
     magnesium oxide (All1(Ba,Mq)CeO19)
                                          145564-56-3P, Calcium magnesium
     silicate ((Ca,Mg)(SiO3)) 146259-99-6P, Strontium boride fluoride oxide
                    186956-28-5P, Aluminum magnesium oxide (All1MgO19)
     (SrB2(F,O)4)
     200212-20-0P, Barium magnesium zinc oxide silicate
     ((Ba, Mg, Zn) 30 (SiO3) 2)
                            225796-98-5P, Aluminum barium magnesium oxide
     (Al2(Ba, Mg)04)
                      230313-54-9P, Gallium yttrium borate ((Ga,Y)(BO3))
     350480-93-2P, Magnesium strontium metaphosphate oxide ((Mg,Sr)2(PO3)20)
     371759-79-4P, Aluminum calcium oxide silicate (Al2Ca2O(SiO3)2)
     371759-81-8P, Aluminum yttrium borate oxide (Al3Y(BO3)3O3)
                                                                  371759-82-9P,
     Aluminum gallium yttrium oxide (Al3Ga2Y2O12)
                                                   944557-34-0P, Aluminum
     barium magnesium oxide (Al13BaMgO23)
                                          944557-35-1P, Aluminum cerium
     magnesium terbium oxide (Al10Ce0.6MgTb0.35019)
                                                      944557-36-2P, Beryllium
     zinc silicate ((Be, Zn)(SiO4))
     RL: IMF (Industrial manufacture); TEM (Technical or engineered material
     use); PREP (Preparation); USES (Uses)
        (synthesis of nanocryst. phosphor particles and the nanocryst. phosphor
        particles and their uses)
     56-81-5, Glycerin, uses
                               64-17-5, Ethanol, uses
                                                        67-56-1, Methanol, uses
     67-68-5, Dimethyl sulfoxide, uses 68-12-2, Dimethylformamide, uses
     75-04-7, Ethylamine, uses 78-50-2, Trioctylphosphine oxide
     Ethylene diamine, uses
                            110-86-1, Pyridine, uses 111-46-6, Diethylene
                   111-86-4, Octylamine 112-40-3, Dodecane
     glycol, uses
                                                                124-07-2.
     Octanoic acid, uses 124-38-9, Carbon dioxide, uses
                                                            143-10-2,
                  151-41-7 4731-53-7, Trioctylphosphine
     Decanethiol
                                                             7446-09-5, Sulfur
     dioxide, uses 7664-41-7, Ammonia, uses
                                                7732-18-5, Water, uses
     39407-03-9, Octyl phosphate 944557-37-3
     RL: NUU (Other use, unclassified); USES (Uses)
        (synthesis of nanocryst. phosphor particles and the nanocryst. phosphor
        particles and their uses)
     78-10-4, Silicon ethoxide
                                5588-84-1
                                             7646-85-7, Zinc chloride,
                 7773-01-5, Manganese chloride
     reactions
                                                 7790-86-5, Cerium chloride
     10025-76-0, Europium chloride 10042-88-3, Terbium chloride
                                                                    10099-58-8,
     Lanthanum trichloride
                             10361-37-2, Barium chloride, reactions
     10361-92-9, Yttrium chloride
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (synthesis of nanocryst. phosphor particles and the nanocryst. phosphor
        particles and their uses)
RE.CNT
              THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
(1) Antonietti, M; ANGEWANDTE CHEMIE 2004, V116, P5096
(2) Deng Tao; US 2005255236 A1 2005
(3) Feldmann, C; ADVANCED FUNCTIONAL MATERIALS 2003, V13(2), P101 CAPLUS
(4) Kompe, K; ANGEWANDTE CHEMIE 2003, V115, P5672
(5) Riwotzki, K; ANGEWANDTE CHEMIE 2001, V113(3), P574
(6) Riwotzki, K; JOURNAL OF PHYSICAL CHEMISTRY B, MATERIALS, SURFACES,
    INTERFACES AND BIOPHYSICAL, WASHINGTON, DC, US 2000, V104, P2824 CAPLUS
ACCESSION NUMBER:
                         2007:816790 CAPLUS
DOCUMENT NUMBER:
                         147:199722
TITLE:
                         Process for the synthesis of nanocrystalline phosphor
                         particles and nanocrystalline phosphor particles
                         obtainable by this process
INVENTOR(S):
                         Feldmann, Claus; Buehler, Gunnar
PATENT ASSIGNEE(S):
                        Universitaet Karlsruhe, Germany
SOURCE:
                         PCT Int. Appl., 34pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                         German
FAMILY ACC. NUM. COUNT:
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ΙT

RE

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PATENT NO.
                                          APPLICATION NO. DATE
                        KIND DATE
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     WO 2007082663
                         A1 20070726 WO 2007-EP175 20070110
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN,
             KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK,
             MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO,
             RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR, TT,
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             CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH,
             GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,
             KG, KZ, MD, RU, TJ, TM
     DE 102006001414
                         A1 20070816
                                          DE 2006-102006001414
PRIORITY APPLN. INFO.:
                                            DE 2006-102006001414A 20060111
     Methods for the targeted synthesis of inorg. phosphor particles which have
     a particle diameter of 1-100 nm, a virtually monodisperse size distribution
     in the range of ±20% and, depending on the class of material selected,
     a quantum yield of \geq 20\% which entail reacting \geq 1 metal
     precursor with ≥1 precipitating reagent at -50° to 200° in a
     first solvent or solvent mixture; thermally treating the resulting mixture at
     150-700^{\circ} for 5 s to 30 min in a second solvent or solvent mixture
     that includes ≥1 ionic liquid; and isolating and/or purifying the
     phosphor particles are described in which the metal precursor is chosen
     from halides, nitrates, sulfates, hydrogen sulfates, acetates,
     acetylacetonates, oxalates, carbonates, hydrogen carbonates, phosphates,
     hydrogen phosphates, dihydrogen phosphates, hydrides, alcoholates, and/or
     metalorg. compds., and the precipitating reagent is chosen from hydroxides,
     phosphates, carbonates, oxalates, sulfates, sulfides, halides, ammonia,
     hydroxylamines, amines, amides, imides, and/or water. Inorg. phosphor
     particles obtainable by this process and also their use for coating or
     embedding in a substrate selected from the group consisting of a paper,
     polymer, glass, metal or ceramic substrate, for example for diagnostics,
     therapy, as contrast agent, as marking, warning feature, for illumination,
     and for data storage and/or data reproduction A particular advantage of the
     phosphor particles which can be obtained according to the invention is
     that the particles have a surface finish which allows agglomerate-free
     redispersion in polar and nonpolar solvents or dispersion media.
     The choice of the dispersion medium allows the physicochem. properties of
     the phosphor particles which can be obtained according to the invention to
     be specifically matched to the requirements of a particular application.
     ANSWER 3 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
L5
AN
     2007:757081 CAPLUS
ΕD
     Entered STN: 12 Jul 2007
ΤI
     Purification and stabilization of colloidal ZnO nanoparticles in
     methanol
ΑU
     Sun, Dazhi; Wong, Minhao; Sun, Luyi; Li, Yuntao; Miyatake, Nobuo; Sue,
     Hung-Jue
CS
     Department of Mechanical Engineering, Polymer Technology Center, Texas A&M
     University, College Station, TX, 77843-3123, USA
     Journal of Sol-Gel Science and Technology (2007), 43(2), 237-243
SO
     CODEN: JSGTEC; ISSN: 0928-0707
PB
     Springer
ĎΤ
     Journal
LA
     English
     66 (Surface Chemistry and Colloids)
CÇ
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Purification and stabilization of colloidal ZnO nanoparticles

synthesized from zinc acetate dihydrate and potassium hydroxide in methanol have been performed. Isopropanol and hexane were utilized to achieve the precipitation-redispersion washing procedure in methanol.

J

AΒ

The results from atomic absorption spectroscopy show that the concentration of K+ cation can be dramatically reduced by washing. X-ray diffraction and transmission electron microscopy results show that zinc layered double hydroxide formed in the ZnO precipitate can be effectively removed through concentrating the unpurified ZnO colloids by solvent evaporation The purified ZnO nanoparticles can be fully redispersed in methanol, but become unstable with time due to the low concentration of acetate in the colloids. Fortunately, the unstable ZnO methanol colloids are found to become stable after addition of a small amount of hexane. The present study is of particular significance for the preparation of purified colloidal ZnO nanoparticles for device fabrication, functional ZnO coatings and polymer nanocomposite applications. ACCESSION NUMBER: 2007:757081 CAPLUS TITLE: Purification and stabilization of colloidal ZnO nanoparticles in methanol AUTHOR(S): Sun, Dazhi; Wong, Minhao; Sun, Luyi; Li, Yuntao; Miyatake, Nobuo; Sue, Hung-Jue CORPORATE SOURCE: Department of Mechanical Engineering, Polymer Technology Center, Texas A&M University, College Station, TX, 77843-3123, USA SOURCE: Journal of Sol-Gel Science and Technology (2007), 43(2), 237-243 CODEN: JSGTEC; ISSN: 0928-0707 PUBLISHER: Springer DOCUMENT TYPE: Journal LANGUAGE: English Purification and stabilization of colloidal ZnO nanoparticles synthesized from zinc acetate dihydrate and potassium hydroxide in methanol have been performed. Isopropanol and hexane were utilized to achieve the precipitation-redispersion washing procedure in methanol. The results from atomic absorption spectroscopy show that the concentration of K+ cation can be dramatically reduced by washing. X-ray diffraction and transmission electron microscopy results show that zinc layered double hydroxide formed in the ZnO precipitate can be effectively removed through concentrating the unpurified ZnO colloids by solvent evaporation The purified ZnO nanoparticles can be fully redispersed in methanol, but become unstable with time due to the low concentration of acetate in the colloids. Fortunately, the unstable ZnO methanol colloids are found to become stable after addition of a small amount of hexane. The present study is of particular significance for the preparation of purified colloidal ZnO nanoparticles for device fabrication, functional ZnO coatings and polymer nanocomposite applications. ANSWER 4 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN L5 ΑN 2007:675926 CAPLUS DN 147:33781 ED Entered STN: 22 Jun 2007 Preparation of metal oxide nanoparticles ΤI ΙN O'Brien, Stephen; Yin, Ming The Trustees of Columbia University In the City of New York, USA PA U.S. Pat. Appl. Publ., 72pp., Cont.-in-part of PCT/2004US/41141. SO CODEN: USXXCO DΤ Patent LA English INCL 423592100; 423632000; 423605000; 423622000 49-3 (Industrial Inorganic Chemicals) Section cross-reference(s): 62, 63, 77 FAN.CNT 2 PATENT NO. KIND DATE APPLICATION NO. ----US 2007140951 A1 20070621 US 2006-451251 20060612 WO 2005060610 A2 20050707 WO 2004-US41141 20041209 WO 2005060610 A3 20050922 PΤ

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W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
             LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
             NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
             TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
         RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK,
             EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT,
             RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML,
             MR, NE, SN, TD, TG
PRAI US 2003-528667P
                         Ρ
                                20031211
     WO 2004-US41141
                                20041209
                          A2
CLASS
 PATENT NO.
                 CLASS PATENT FAMILY CLASSIFICATION CODES
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 US 2007140951
                 INCL
                        423592100; 423632000; 423605000; 423622000
                 IPCI
                        C01B0013-14 [I,A]
                 NCL
                        423/592.100; 423/632.000; 423/605.000; 423/622.000
WO 2005060610
                 IPCI
                        A61K [ICM, 7]
                 IPCR
                        C01F0017-00 [I,C*]; C01F0017-00 [I,A]; C01G0003-00
                        [I,C*]; C01G0003-00 [I,A]; C01G0009-00 [I,C*];
                        C01G0009-00 [I,A]; C01G0023-00 [I,C*]; C01G0023-00
                        [I,A]; C01G0025-00 [I,C*]; C01G0025-00 [I,A];
                        C01G0027-00 [I,C*]; C01G0027-00 [I,A]; C01G0031-00
                        [I,C*]; C01G0031-00 [I,A]; C01G0037-00 [I,C*];
                        C01G0037-00 [I,A]; C01G0039-00 [I,C*]; C01G0039-00
                        [I,A]; C01G0045-00 [I,C*]; C01G0045-00 [I,A];
                        C01G0049-00 [I,C*]; C01G0049-00 [I,A]; C01G0051-00
                        [I,C*]; C01G0051-00 [I,A]; C01G0055-00 [I,C*];
                        C01G0055-00 [I,A]
AΒ
     Metal oxide nanoparticles are prepared by (a) mixing a metal acetate
     precursor with a nonaq. organic solvent containing at least one organic
stabilizing
     ligand to form a reaction mixture; (b) subjecting the mixture of step (a) to a
     temperature of 250-400° for a time sufficient to allow formation of metal
     oxide nanoparticles and decomposition of the metal acetate; and (c) extracting
the
    metal oxide nanoparticles into a hydrocarbon solvent at a temperature lower
than
     the temperature of step (b) by precipitation with a flocculating agent.
solvent is
     trioctylamine and the organic stabilizing ligand is oleic acid.
    Nanoparticles produced can be oxides of Zn, Fe, Mn, Co, Ru, Cu, Sc, Ti, V,
    Cr, Mo, Y, Zr, Hf, or Ni. The hydrocarbon solvent used in step (c) can be
    pentane, hexane, heptane, octane, or dodecane. The flocculating agent
    comprises a polar organic solvent, such as methanol, ethanol, propanol, or
    butanol. Preferably, the nanoparticles are extracted into hexane solvent by
    precipitation with ethanol, followed by centrifugation and redispersion
    in hydrocarbon solvent. The extracted nanoparticles are stable, monodisperse,
    and have uniform size. The extracted metal oxide nanoparticles may be
    subjected to oxidation to obtain further oxidized metal oxide nanoparticles.
    The resulting nanoparticles can exhibit magnetic and/or optical
    properties. The nanoparticles are well suited for use in a variety of
    industrial applications, including cosmetic and pharmaceutical
    formulations and compns. The cosmetic formulation can be makeups, topical
    skin care products, soaps, powders, lotions, creams, ointments, sunblocks,
    sunscreens, conditioners, shampoos, fragrances, deodorants, deodorizers,
    hair colors, or hair dyes.
ST
    metal oxide nanoparticle prepn trioctylamine oleic acid
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(nanoparticles; preparation of metal oxide nanoparticles)
IT Magnetic materials
 Nanoparticles
 Optical filters

Drug delivery systems

TT

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Particle size
     Sunscreens
         (preparation of metal oxide nanoparticles)
ΙT
     Oxides (inorganic), preparation
     RL: IMF (Industrial manufacture); NUU (Other use, unclassified); PEP
     (Physical, engineering or chemical process); PRP (Properties); PREP
     (Preparation); PROC (Process); USES (Uses)
        (preparation of metal oxide nanoparticles)
     1344-43-0P, Manganese oxide (MnO), preparation
ΙT
     RL: IMF (Industrial manufacture); PEP (Physical, engineering or chemical
     process); PRP (Properties); PREP (Preparation); PROC (Process)
        (nanoparticles; preparation of metal oxide nanoparticles)
ΙT
     1184-78-7, Trimethylamine N-oxide
     RL: PEP (Physical, engineering or chemical process); PROC (Process)
        (oxidant; preparation of metal oxide nanoparticles)
     1309-37-1P, Iron oxide (Fe2O3), preparation
TΤ
                                                   1314-13-2P, Zinc
     oxide, preparation
                          1317-35-7P, Manganese oxide (Mn3O4)
     1317-38-0P, Copper oxide (CuO), preparation 1317-39-1P, Copper oxide,
                   1317-61-9P, Iron oxide (Fe3O4), preparation 1345-25-1P,
     preparation
     Iron oxide (FeO), preparation
     RL: IMF (Industrial manufacture); PEP (Physical, engineering or chemical
     process); PRP (Properties); PREP (Preparation); PROC (Process)
        (preparation of metal oxide nanoparticles)
ΙT
     112-80-1, Oleic acid, uses
                                  1116-76-3, Trioctylamine
     RL: NUU (Other use, unclassified); USES (Uses)
        (preparation of metal oxide nanoparticles)
ΙT
     142-71-2, Copper II acetate
                                  638-38-0, Manganese II acetate
                                                                      3094-87-9,
     Iron II acetate
     RL: PEP (Physical, engineering or chemical process); RCT (Reactant); PROC
     (Process); RACT (Reactant or reagent)
        (preparation of metal oxide nanoparticles)
                         2007:675926 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         147:33781
TITLE:
                         Preparation of metal oxide nanoparticles
INVENTOR(S):
                         O'Brien, Stephen; Yin, Ming
PATENT ASSIGNEE(S):
                         The Trustees of Columbia University In the City of New
                         York, USA
SOURCE:
                         U.S. Pat. Appl. Publ., 72pp., Cont.-in-part of
                         PCT/2004US/41141.
                         CODEN: USXXCO
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT: 2
PATENT INFORMATION:
     PATENT NO.
                        KIND
                                DATE
                                           APPLICATION NO.
                                                                    DATE
                         ____
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     US 2007140951
                         A1
                                20070621
                                             US 2006-451251
                                                                     20060612
     WO 2005060610
                         A2
                                 20050707
                                            WO 2004-US41141
                                                                    20041209
                         A3
     WO 2005060610
                               20050922
            AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
             CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
             GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
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GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW, RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG

PRIORITY APPLN. INFO.:

US 2003-528667P P 20031211

WO 2004-US41141 A2 20041209

AB Metal oxide nanoparticles are prepared by (a) mixing a metal acetate precursor with a nonaq. organic solvent containing at least one organic stabilizing
```

ligand to form a reaction mixture; (b) subjecting the mixture of step (a) to a temperature of $250\text{--}400^{\circ}$ for a time sufficient to allow formation of metal oxide nanoparticles and decomposition of the metal acetate; and (c) extracting

the

metal oxide nanoparticles into a hydrocarbon solvent at a temperature lower than

the temperature of step (b) by precipitation with a flocculating agent. The solvent is

trioctylamine and the organic stabilizing ligand is oleic acid. Nanoparticles produced can be oxides of Zn, Fe, Mn, Co, Ru, Cu, Sc, Ti, V, Cr, Mo, Y, Zr, Hf, or Ni. The hydrocarbon solvent used in step (c) can be pentane, hexane, heptane, octane, or dodecane. The flocculating agent comprises a polar organic solvent, such as methanol, ethanol, propanol, or butanol. Preferably, the nanoparticles are extracted into hexane solvent by precipitation with ethanol, followed by centrifugation and redispersion in hydrocarbon solvent. The extracted nanoparticles are stable, monodisperse, and have uniform size. The extracted metal oxide nanoparticles may be subjected to oxidation to obtain further oxidized metal oxide nanoparticles. The resulting nanoparticles can exhibit magnetic and/or optical properties. The nanoparticles are well suited for use in a variety of industrial applications, including cosmetic and pharmaceutical formulations and compns. The cosmetic formulation can be makeups, topical skin care products, soaps, powders, lotions, creams, ointments, sunblocks, sunscreens, conditioners, shampoos, fragrances, deodorants, deodorizers, hair colors, or hair dyes.

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ANSWER 5 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
L5
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AN 2006:982553 CAPLUS

DN 145:340496

ED Entered STN: 22 Sep 2006

ΤI Reactivity and concentration in size and shape control of metal oxide nanocrystals in synthesis by reaction of metal fatty acid salt, fatty acid and hydrocarbon solvent and pyrolysis of product

Peng, Xiaogang; Chen, Yonfen; Jana, Nikhil; Narayanaswamy, Arun IN

PΑ The Board of Trustees of the University of Arkansas, USA SO

PCT Int. Appl., 37pp.

CODEN: PIXXD2

DT Patent

LA English

CC 57-2 (Ceramics)

Section cross-reference(s): 78

FAN.CNT 1

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PI		2006098756 2006098756					20060921 20070426		WO 2005-US29072					20050816				
		W:	CN, GE, LC, NG, SL, ZA,	CO, GH, LK, NI, SM, ZM,	CR, GM, LR, NO, SY, ZW	CU, HR, LS, NZ, TJ,	CZ, HU, LT, OM, TM,	AU, DE, ID, LU, PG, TN,	DK, IL, LV, PH, TR,	DM, IN, MA, PL, TT,	DZ, IS, MD, PT, TZ,	EC, JP, MG, RO, UA,	EE, KE, MK, RU, UG,	EG, KG, MN, SC, US,	ES, KM, MW, SD, UZ,	FI, KP, MX, SE, VC,	GB, KR, MZ, SG, VN,	GD, KZ, NA, SK, YU,
	nc.		IS, CF, GM, KG,	IT, CG, KE, KZ,	LT, CI, LS, MD,	LU, CM, MW, RU,	LV, GA, MZ, TJ,	MC, GN, NA, TM,	NL, GQ, SD, AP,	PL, GW, SL, EA,	PT, ML, SZ, EP,	RO, MR, TZ, OA	SE, NE, UG,	SI, SN, ZM,	SK, TD, ZW,	TR, TG, AM,	BF, BW, AZ,	BJ, GH, BY,
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DDAT		2006						2006		1	US 2	005-2	2047	66		20	0050	316
PRAI		2004																
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WO 2006098756
                IPCI
                         H01L0021-00 [I,A]; C01G0009-00 [I,C]; C01G0009-02
                         [I,A]; B32B0005-16 [I,C]; B32B0005-16 [I,A];
                         C01G0037-00 [I,C]; C01G0037-02 [I,A]; C01G0049-02
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                         C01G0053-04 [I,A]
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 US 7160525
                  IPCI
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                         [I,A]; C01G0055-00 [I,A]
                  IPCR
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                         [I,C]; C01G0007-00 [I,A]; C01G0051-00 [I,C];
                         C01G0051-00 [I,A]; C01G0055-00 [I,C]; C01G0055-00 [I,A]
                  NCL
                         423/001.000; 423/022.000; 423/023.000; 423/024.000;
                         423/138.000; 423/139.000
                  ECLA
                         MO1P; MO1P; MO1P; MO1P; M22B; M22B; Y01N
 US 2006211152
                  IPCI
                         H01L0021-00 [I,A]
                  IPCR
                         H01L0021-00 [I,C]; H01L0021-00 [I,A]
                  NCL
                         438/003.000; 977/963.000
AΒ
     A general, reproducible, and simple synthetic method that employs readily
     available chems. permits control of the size, shape, and size distribution
     of metal oxide nanocrystals. The synthesis entails reacting a metal fatty acid salt, the corresponding fatty acid, and a hydrocarbon solvent, with
     the reaction product being pyrolyzed to the metal oxide. Nearly
     monodisperse oxide nanocrystals of Fe3O4, Cr2O3, MnO, Co3O4, NiO,
     ZnO, SnO2, and In2O3, in a large size range (3-50 nm), are
     described. Size and shape control of the nanocrystals is achieved by
     varying the reactivity and concentration of the precursors.
ST
     metal oxide nanocrystal synthesis size shape control reactivity concn;
     fatty acid salt hydrocarbon solvent metal oxide nanocrystal synthesis
ΙT
     Powders
        (ceramic; size and shape control of metal oxide nanocrystals in
        synthesis by reaction of metal fatty acid salt, fatty acid and
        hydrocarbon solvent and pyrolysis)
ΙT
     Nanocrystals
        (oxide; size and shape control of metal oxide nanocrystals in synthesis
        by reaction of metal fatty acid salt, fatty acid and hydrocarbon
        solvent and pyrolysis)
IT
     Ceramics
        (powders; size and shape control of metal oxide nanocrystals in
        synthesis by reaction of metal fatty acid salt, fatty acid and
        hydrocarbon solvent and pyrolysis)
ΙT
     Fatty acids, reactions
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (salts; size and shape control of metal oxide nanocrystals in synthesis
        by reaction of metal fatty acid salt, fatty acid and hydrocarbon
        solvent and pyrolysis)
IT
     Concentration (condition)
     Crystal morphology
     Grain size
     Particle size
     Reactivity (chemical)
     Solvents
     Thermal decomposition
        (size and shape control of metal oxide nanocrystals in synthesis by
        reaction of metal fatty acid salt, fatty acid and hydrocarbon solvent
        and pyrolysis)
ΙT
     Fatty acids, reactions
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (size and shape control of metal oxide nanocrystals in synthesis by
        reaction of metal fatty acid salt, fatty acid and hydrocarbon solvent
        and pyrolysis)
ΙT
     Hydrocarbons, reactions
     RL: NUU (Other use, unclassified); RCT (Reactant); RACT (Reactant or
     reagent); USES (Uses)
        (solvents; size and shape control of metal oxide nanocrystals in
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synthesis by reaction of metal fatty acid salt, fatty acid and
        hydrocarbon solvent and pyrolysis)
     1308-06-1P, Cobalt oxide (Co304)
TΤ
                                         1308-38-9P, Chromium oxide (Cr203),
                   1312-43-2P, Indium oxide (In2O3)
     preparation
                                                       1313-99-1P, Nickel oxide
     (NiO), preparation 1314-13-2P, Zinc oxide (
     ZnO), preparation 1317-61-9P, Iron oxide (Fe3O4), preparation
     1344-43-0P, Manganous oxide, preparation
                                                 18282-10-5P, Tin oxide (SnO2)
     RL: SPN (Synthetic preparation); PREP (Preparation)
        (nanocrystals; size and shape control of metal oxide nanocrystals in
        synthesis by reaction of metal fatty acid salt, fatty acid and
        hydrocarbon solvent and pyrolysis)
ΙT
     108-88-3, Toluene, uses
     RL: NUU (Other use, unclassified); USES (Uses)
        (redispersion in; size and shape control of metal oxide
        nanocrystals in synthesis by reaction of metal fatty acid salt, fatty acid and hydrocarbon solvent and pyrolysis)
IT
     57-11-4, Stearic acid, reactions 112-80-1, Oleic acid, reactions
     544-63-8, Myristic acid, reactions 557-05-1, Zinc stearate 2800-96-6,
     Tin(IV) acetate 13586-84-0, Cobalt stearate
                                                     14448-69-2, Nickel
     stearate
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (size and shape control of metal oxide nanocrystals in synthesis by
        reaction of metal fatty acid salt, fatty acid and hydrocarbon solvent
        and pyrolysis)
     1120-45-2P, Ferric oleate
ΙT
                                2980-59-8P, Iron(II) stearate
                                                                  15114-27-9P,
     Ferrous oleate 65526-79-6P, Indium stearate
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (size and shape control of metal oxide nanocrystals in synthesis by
        reaction of metal fatty acid salt, fatty acid and hydrocarbon solvent
        and pyrolysis)
ΤТ
     112-95-8, n-Eicosane
                             646-31-1, Tetracosane
                                                     27070-58-2, Octadecene
     RL: NUU (Other use, unclassified); RCT (Reactant); RACT (Reactant or
     reagent); USES (Uses)
        (solvent; size and shape control of metal oxide nanocrystals in
        synthesis by reaction of metal fatty acid salt, fatty acid and
        hydrocarbon solvent and pyrolysis)
ACCESSION NUMBER:
                         2006:982553 CAPLUS
DOCUMENT NUMBER:
                         145:340496
TITLE:
                         Reactivity and concentration in size and shape control
                         of metal oxide nanocrystals in synthesis by reaction
                         of metal fatty acid salt, fatty acid and hydrocarbon
                         solvent and pyrolysis of product
INVENTOR(S):
                         Peng, Xiaogang; Chen, Yonfen; Jana, Nikhil;
                         Narayanaswamy, Arun
PATENT ASSIGNEE(S):
                         The Board of Trustees of the University of Arkansas,
                         USA
SOURCE:
                         PCT Int. Appl., 37pp.
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
     PATENT NO
                         KTND
                                             A DOLLCATION NO
                                שאתב
     ___
     WO
     WO
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LENI	NO.			KIN	D	DATE			APPL.	ICAT.	TON I	NO.		D2	ATE		
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2006	0987	56		A2		2006	0921	1	WO 2	005-1	US29	072		21	00508	816	
2006	0987	56		A3		2007	0426										
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						DE,											
	GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KP,	KR,	KZ,	
						LU,											
						PG,											
						TN,											
		ZM,												-	•	•	

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RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
              IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ,
             CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AP, EA, EP, OA
     US 7160525
                         B1 20070109 US 2004-965685
A1 20060921 US 2005-204766
                                                                     20041014
     US 2006211152
                                              US 2005-204766
                                                                     20050816
PRIORITY APPLN. INFO.:
                                              US 2004-601778P
                                                                 P 20040816
                                              US 2004-965685
                                                               A 20041014
P 20031014
                                              US 2003-511183P
     A general, reproducible, and simple synthetic method that employs readily
AB
     available chems. permits control of the size, shape, and size distribution
     of metal oxide nanocrystals. The synthesis entails reacting a metal fatty acid salt, the corresponding fatty acid, and a hydrocarbon solvent, with
     the reaction product being pyrolyzed to the metal oxide. Nearly
     monodisperse oxide nanocrystals of Fe304, Cr203, MnO, Co304, NiO,
     ZnO, SnO2, and In2O3, in a large size range (3-50 nm), are
     described. Size and shape control of the nanocrystals is achieved by
     varying the reactivity and concentration of the precursors.
L5
     ANSWER 6 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
ΑN
     2006:68445 CAPLUS
DN
     144:214450
ED
     Entered STN: 25 Jan 2006
     Decorative coating with air purifying, antibacterial and humidity
ΤI
     adjusting effects for internal wall
     Ji, Zhijiang; Zhang, Liansong; Wang, Jing; Wang, Jimei; Wang, Xiaoyan; Lu,
IN
     Rongchao
     China Building Materials Academy, Peop. Rep. China
PA
SO
     Faming Zhuanli Shenqing Gongkai Shuomingshu, 11 pp.
     CODEN: CNXXEV
DT
     Patent
LA
     Chinese
     ICM C09D131-04
IC
     ICS C09D129-04; C09D005-03
     42-10 (Coatings, Inks, and Related Products)
FAN.CNT 1
     PATENT NO.
                         KIND
                                 DATE
                                            APPLICATION NO. DATE
                        ----
                                 _____
                                            -----
    CN 1632010
PΙ
                         Α
                                 20050629
                                            CN 2004-10088977
                                                                   20041123
PRAI CN 2004-10088977
                                 20041123
CLASS
 PATENT NO.
             CLASS PATENT FAMILY CLASSIFICATION CODES
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 CN 1632010
                ICM
                         C09D131-04
                 ICS
                         C09D129-04; C09D005-03
                 IPCI
                         C09D0131-04 [ICM, 7]; C09D0131-00 [ICM, 7, C*];
                         C09D0129-04 [ICS,7]; C09D0129-00 [ICS,7,C*];
                         C09D0005-03 [ICS,7]
                 IPCR
                         C09D0005-03 [I,C*]; C09D0005-03 [I,A]; C09D0129-00
                         [I,C*]; C09D0129-04 [I,A]; C09D0131-00 [I,C*];
                         C09D0131-04 [I,A]
AB
     The title decorative coating contains nanomaterial 2-10, moisture
     absorbing material 10-40, inorg. filler 50-80, and redispersible
     powder 2-5 wt%. With the synergetic effect of various mineral materials
     such as inorg. mineral material, natural microporous or layered material,
     nanomaterial and redispersible powder, the coating has special
     functions including capability of releasing neg. ions. It is also
     environment-friendly as it contains no VOC.
ST
     decorative coating nanomaterial moisture absorbing inorg filler
     redispersible powder
IT
     Absorbents
     Cement
     Fillers
     Nanoparticles
```

Powders (coating containing; preparation of decorative coating with air purifying, antibacterial and humidity adjusting effects for internal wall) IT Bentonite, uses Diatomite Kaolin, uses Tourmaline-group minerals Zeolites (synthetic), uses RL: PRP (Properties); TEM (Technical or engineered material use); USES (Uses) (coating containing; preparation of decorative coating with air purifying, antibacterial and humidity adjusting effects for internal wall) ΙT Antibacterial agents Coating materials (preparation of decorative coating with air purifying, antibacterial and humidity adjusting effects for internal wall) ΙT 87714-59-8, DM 200 RL: PRP (Properties); TEM (Technical or engineered material use); USES (Uses) (DM 200, coating containing; preparation of decorative coating with air purifying, antibacterial and humidity adjusting effects for internal wall) 108-05-4D, Vinyl acetate, polymers with vinyl versatate 1305-62-0, ΤТ Calcium hydroxide, uses 1309-37-1, Iron oxide, uses 1314-13-2, Zinc oxide, uses 1314-35-8, Tungsten oxide, uses 1332-29-2, Tin oxide 7439-91-0D, Lanthanum, salts or oxides 7440-00-8D, Neodymium, salts or oxides 7440-45-1D, Cerium, 7440-45-1D, Cerium, salts or oxides 7631-86-9, Silica, uses 9002-89-5, Polyvinyl alcohol 9004-62-0, Hydroxyethyl cellulose 9032-42-2, Hydroxyethyl methyl cellulose 12174-11-7, Attapulgite 13397-24-5, Gypsum, uses 13463-67-7, Titania, uses 14807-96-6, Talc, uses 26221-27-2, Ethylene-vinyl acetate-vinyl alcohol copolymer 52383-89-8, Vinnapas RI 63800-37-3, Sepiolite 408325-37-1, Tylovis SE 7 875690-83-8, 551Z Rovace DP 87 RL: PRP (Properties); TEM (Technical or engineered material use); USES (Uses) (coating containing; preparation of decorative coating with air purifying, antibacterial and humidity adjusting effects for internal wall) 471-34-1, Calcium carbonate, uses IT RL: PRP (Properties); TEM (Technical or engineered material use); USES (high-d., coating containing; preparation of decorative coating with air purifying, antibacterial and humidity adjusting effects for internal wall) ΙT 9005-25-8D, Starch, ether derivs. RL: PRP (Properties); TEM (Technical or engineered material use); USES (preparation of decorative coating with air purifying, antibacterial and humidity adjusting effects for internal wall) ACCESSION NUMBER: 2006:68445 CAPLUS DOCUMENT NUMBER: 144:214450 Decorative coating with air purifying, antibacterial TITLE: and humidity adjusting effects for internal wall INVENTOR(S): Ji, Zhijiang; Zhang, Liansong; Wang, Jing; Wang, Jimei; Wang, Xiaoyan; Lu, Rongchao PATENT ASSIGNEE(S): China Building Materials Academy, Peop. Rep. China SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 11 pp. CODEN: CNXXEV DOCUMENT TYPE: Patent LANGUAGE: Chinese FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

APPLICATION NO.

DATE

PATENT NO.

KIND

DATE

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CN 1632010
                         A
                                20050629 CN 2004-10088977
                                            CN 2004-10088977
PRIORITY APPLN. INFO.:
     The title decorative coating contains nanomaterial 2-10, moisture
     absorbing material 10-40, inorg. filler 50-80, and redispersible
     powder 2-5 wt%. With the synergetic effect of various mineral materials
     such as inorg. mineral material, natural microporous or layered material,
     nanomaterial and redispersible powder, the coating has special
     functions including capability of releasing neg. ions. It is also
     environment-friendly as it contains no VOC.
L5
     ANSWER 7 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
AN
     2004:1127444 CAPLUS
DN
     142:58371
ΕD
     Entered STN: 24 Dec 2004
     Nanoparticulate redispersible zinc oxide
ΤI
     powder iii
     Kliss, Rainer; Kropf, Christian; Hahn, Horst; Bergmann, Soeren; Umbreit,
ΤN
     Christian; Peschke, Joerg
PΑ
     Sustech GmbH & Co. Kg, Germany
     PCT Int. Appl., 52 pp.
SO
     CODEN: PIXXD2
DT
     Patent
LA
     German
IC
     ICM C09C001-04
     ICS C01G009-02; C08K003-22; C08K009-04; A61K007-48; A61K007-42
     42-10 (Coatings, Inks, and Related Products)
CC
FAN.CNT 1
                               DATE APPLICATION NO.
     PATENT NO.
                       KIND DATE
                                                              DATE
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                         ____
                                            ______
    WO 2004111136 A1 20041223 WO 2004-EP6463 20040616
PΤ
        RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,
             SN, TD, TG
     DE 10327728
                         A1
                               20050127 DE 2003-10327728 20030618
PRAI DE 2003-10327728
                        Α
                                20030618
CLASS
PATENT NO.
                CLASS PATENT FAMILY CLASSIFICATION CODES
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WO 2004111136
                 ICM
                        C09C001-04
                 ICS
                        C01G009-02; C08K003-22; C08K009-04; A61K007-48;
                        A61K007-42
                 IPCI
                        C09C0001-04 [ICM,7]; C01G0009-02 [ICS,7]; C01G0009-00
                        [ICS,7,C*]; C08K0003-22 [ICS,7]; C08K0003-00
                        [ICS,7,C*]; C08K0009-04 [ICS,7]; C08K0009-00
                        [ICS,7,C*]; A61K0007-48 [ICS,7]; A61K0007-42 [ICS,7]
                        A61K0008-19 [I,C*]; A61K0008-27 [I,A]; A61Q0017-04
                 IPCR
                        [I,C*]; A61Q0017-04 [I,A]; A61Q0019-00 [I,C*];
                        A61Q0019-00 [I,A]; C09C0001-04 [I,C*]; C09C0001-04
                        [I,A]
                 ECLA
                        A61K008/27; A61Q017/04; A61Q019/00; C09C001/04B
DE 10327728
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                        C01G0009-02 [ICM, 7]; C01G0009-00 [ICM, 7, C*];
                        A61K0007-48 [ICS, 7]
                 TPCR
                        A61K0008-19 [I,C*]; A61K0008-27 [I,A]; A61Q0017-04
                        [I,C*]; A61Q0017-04 [I,A]; A61Q0019-00 [I,C*];
                        A61Q0019-00 [I,A]; C09C0001-04 [I,C*]; C09C0001-04
                        [I,A]
                 ECLA
                        A61K008/27; A61Q017/04; A61Q019/00; C09C001/04B
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Surface-modified redispersible zinc oxide
AΒ
     nanoparticles [prepared by coating of ZnO with organic acids
     R1CH: CHR2O(CH2CH2O) nCH2COOH (R1 and R2 = H or C1-30 alkyl, n = 1 - 30)
     for manufacture of long-term stable dispersions in organic solvents are useful
in
     cosmetics, for UV-protection and as antimicrobial agents. Thus, adding a
     solution 6.2 g of ethercarboxylic acid in toluene to a suspension of
     ZnO particles having diameter 6 nm (prepared by hydrolysis of ZnCl2 in
     MeOH with aqueous solution of NaOH and washing in MeOH and toluene followed by
     centrifugation), mixing 3 h at elevated temperature, cooling to room
temperature,
     centrifuging and drying the rest at 50° gave a powder forming a
     long-term stable transparent dispersion in toluene having particle size 23
ST
     surface modified redispersible zinc oxide
     nanoparticle coated org acid; surface modified redispersible
     zinc oxide nanoparticle cosmetic antimicrobial use
ΙT
     Coating materials
        (UV-absorbing; surface-modified redispersible zinc
        oxide nanoparticles prepared by coating of ZnO with
        unsat. organic acids for manufacture of stable dispersions useful for
        UV-protection)
IT
     Antimicrobial agents
        (surface-modified redispersible zinc oxide
        nanoparticles prepared by coating of ZnO with unsat. organic acids
        for manufacture of stable dispersions useful as antimicrobial agents)
IT
     Cosmetics
        (surface-modified redispersible zinc oxide
        nanoparticles prepared by coating of ZnO with unsat. organic acids
        for manufacture of stable dispersions useful in cosmetics)
     Carboxylic acids, processes
ΙT
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (unsatd.; surface-modified redispersible zinc
        oxide nanoparticles prepared by coating of ZnO with
        unsatd. organic acids)
ΙT
     57635-48-0
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (organic acids; surface-modified redispersible zinc
        oxide nanoparticles prepared by coating of ZnO with
        unsat. organic acids)
     7646-85-7, Zinc chloride (ZnCl2), reactions
IT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (surface-modified redispersible zinc oxide
        nanoparticles prepared by coating of ZnO with unsat. organic
        acids)
ΙT
     1314-13-2, Zinc oxide (ZnO), uses
     RL: TEM (Technical or engineered material use); USES (Uses)
        (surface-modified redispersible zinc oxide
        nanoparticles prepared by coating of ZnO with unsat. organic
        acids)
              THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE.CNT
(1) Bayer Ag; EP 1157064 A 2001 CAPLUS
(2) Cognis Deutschland Gmbh; WO 0249559 A 2002
(3) Hell; WO 03053398 A 2003 CAPLUS
(4) Johnson & Johnson Consumer; EP 0433086 A 1991 CAPLUS
(5) Pq Corp; WO 9524359 A 1995 CAPLUS
ACCESSION NUMBER:
                         2004:1127444 CAPLUS
DOCUMENT NUMBER:
                         142:58371
TITLE:
                         Nanoparticulate redispersible zinc
                         oxide powder iii
INVENTOR(S):
                         Kliss, Rainer; Kropf, Christian; Hahn, Horst;
                         Bergmann, Soeren; Umbreit, Christian; Peschke, Joerg
```

PATENT ASSIGNEE(S):

Sustech GmbH & Co. Kg, Germany

SOURCE:

PCT Int. Appl., 52 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATE	NT INFORMATION:								
		KIND	DATE	APPLICATION NO.	DATE				
	WO 2004111136 W: AE, AG, AL, CN, CO, CR, GH, GM, HR, LR, LS, LT, NZ, OM, PG, TM, TN, TR, RW: BW, GH, GM, AZ, BY, KG, EE, ES, FI, SI, SK, TR, SN, TD, TG	A1 AM, AT CU, CZ HU, ID LU, LV PH, PL TT, TZ KE, LS KZ, MD FR, GB	20041223 , AU, AZ, BA , DK, DM, DZ , IL, IN, IS , MA, MD, MG , PT, RO, RU , UA, UG, US , MW, MZ, NA , RU, TJ, TM , GR, HU, IE	WO 2004-EP6463 A, BB, BG, BR, BW, BY E, EC, EE, EG, ES, FI G, JP, KE, KG, KP, KR G, MK, MN, MW, MX, MZ J, SC, SD, SE, SG, SK G, UZ, VC, VN, YU, ZA A, SD, SL, SZ, TZ, UG J, AT, BE, BG, CH, CY L, IT, LU, MC, NL, PL L, CM, GA, GN, GQ, GW	, BZ, CA, CH, , GB, GD, GE, , KZ, LC, LK, , NA, NI, NO, , SL, SY, TJ, , ZM, ZW , ZM, ZW, AM, , CZ, DE, DK, , PT, RO, SE,				
DD T O	DE 10327728 RITY APPLN. INFO.:	A1	20050127	DE 2003-10327728					
AB	Surface-modified re	dispers	ible zinc ox	DE 2003-10327728	A 20030618				
	nanoparticles [prep	ared by	coating of	ZnO with organic acid	ds				
	for manufacture of	nCH2COOI	H (R1 and R2 rm stable di	c = H or C1-30 alkyl, spersions in organic	n = 1 - 30)] solvents are useful				
in	cosmetics for IV-n	rotocti	on and as an	timicrobial agents.	Thus adding a				
tempe	solution 6.2 g of e ZnO particles havin MeOH with aqueous s	thercar g diame olution	boxylic acid ter 6 nm (pr of NaOH and	l in toluene to a suspension of the state of	pension of of ZnCl2 in toluene followed by				
•	centrifuging and dr	ying the anspare	e rest at 50 nt dispersio	° gave a powder form n in toluene having p	ing a particle size 23				
L5 AN DN ED TI	ANSWER 8 OF 26 CAP 2004:509968 CAPLUS 141:59224 Entered STN: 24 Ju Redispersible nanop powder with surface conducting surfaces	n 2004 articula coating	ate zinc oxi		lectrical				
IN		iss, Ra:	iner; Hahn,	Horst; Kropf, Christ	ian; Berber,				
PA SO	Sustech GmbH & Co. Ger. Offen., 19 pp.								
DT	CODEN: GWXXBX Patent								
LA	German								
IC	ICM C01G009-02								
ICS A61K007-42 CC 62-4 (Essential Oils and Cosmetics)									
FAN.C	Section cross-refer	ence(s):	: 49						
	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE				
PI	DE 10257388 WO 2004052327	A1 A2	20040624 20040624	DE 2002-10257388 WO 2003-EP13286	20021206 20031126				
	WO 2004052327 W: AU, BR, BY, SG, UA, US,			, IN, JP, KR, MX, NO,	NZ, PL, RU,				

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RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
             IT, LU, MC, NL, PT, RO, SE, SI, SK, TR
     AU 2003288172
                                20040630 AU 2003-288172
                          A1
                                                                    20031126
     EP 1567121
                          Α2
                                20050831
                                           EP 2003-780059
                                                                   20031126
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK
PRAI DE 2002-10257388
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     WO 2003-EP13286
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                                20031126
CLASS
 PATENT NO.
                 CLASS PATENT FAMILY CLASSIFICATION CODES
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                        C01G009-02
 DE 10257388
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                 ICS
                        A61K007-42
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                        C01G0009-02 [ICM,7]; C01G0009-00 [ICM,7,C*];
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                        A61K0033-30 [I,A]; A61Q0019-00 [I,C*]; A61Q0019-00
                        [I,A]; A61Q0019-06 [I,C*]; A61Q0019-06 [I,A];
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 WO 2004052327
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                        A61K0007-42 [ICM,7]; A61K0033-30 [ICS,7]; C01G0009-02
                        [ICS,7]; C01G0009-00 [ICS,7,C*]; C08K0003-22 [ICS,7];
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                        A61K0033-30 [I,A]; A61Q0019-00 [I,C*]; A61Q0019-00
                        [I,A]; A61Q0019-06 [I,C*]; A61Q0019-06 [I,A];
                        C01G0009-00 [I,C*]; C01G0009-02 [I,A]; C09C0001-04
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                        A61K033/30; C01G009/02; C09C001/04B; A01N057/16+M;
                        A61K008/91; A61Q019/00; A61Q019/06
 AU 2003288172
                 IPCI
                        A61K0007-42 [ICM, 7]; C09C0001-04 [ICS, 7]; A61K0033-30
                        [ICS,7]; C01G0009-02 [ICS,7]; C01G0009-00 [ICS,7,C*];
                        C08K0003-22 [ICS,7]; C08K0003-00 [ICS,7,C*];
                        C08K0009-04 [ICS,7]; C08K0009-00 [ICS,7,C*]
                        A01N0057-00 [I,C*]; A01N0057-16 [I,A]; A61K0008-72
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                        [I,C*]; A61K0008-91 [I,A]; A61K0033-30 [I,C*];
                        A61K0033-30 [I,A]; A61Q0019-00 [I,C*]; A61Q0019-00
                        [I,A]; A61Q0019-06 [I,C*]; A61Q0019-06 [I,A];
                        C01G0009-00 [I,C*]; C01G0009-02 [I,A]; C09C0001-04
                        [I,C*]; C09C0001-04 [I,A]
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                 IPCI
                        A61K0007-42 [ICM,7]; A61K0033-30 [ICS,7]; C01G0009-02
                        [ICS,7]; C01G0009-00 [ICS,7,C*]; C08K0003-22 [ICS,7];
                        C08K0003-00 [ICS,7,C*]; C08K0009-04 [ICS,7];
                        C08K0009-00 [ICS,7,C*]; C09C0001-04 [ICS,7]
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                        [I,C*]; A61K0008-91 [I,A]; A61K0033-30 [I,C*];
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                        [I,A]; A61Q0019-06 [I,C*]; A61Q0019-06 [I,A];
                        C01G0009-00 [I,C*]; C01G0009-02 [I,A]; C09C0001-04
                        [I,C*]; C09C0001-04 [I,A]
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                        A01N057/16+M; A61K008/91; A61K033/30; A610019/00;
                        A61Q019/06; C01G009/02; C09C001/04B; M01P; M01P; M01P;
                        M01P; Y01N
OS
    MARPAT 141:59224
     The invention concerns nanoparticular zinc oxide that
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AB is surface-modified with an organic acid, e.g. isostearic acid or polyglycolic acid derivs. The surface-treated zinc oxide can be redispersed in polar and apolar organic solvents. Particles for elec. conducting surfaces and cosmetics are prepared Thus zinc oxide was precipitated by dissolving 1110 g zinc chloride

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in 11.1 L methanol and adding 651 g sodium hydroxide pellets under
     stirring and maintaining room temperature The NaCl/ZnO mixture was
     separated, followed by washing and resuspension in 11 L THF. 385 G isostearic
     acid were added; after 3 h stirring was stopped; sodium chloride settled
     and coated zinc oxide was contained in the clear
     supernatant. The clear supernatant was separated, the solvent evaporated and
the
     raw product reconstructed in toluene; the residual sodium chloride was
     removed by centrifugation. 4L ethanol (96%) were added to the centrifuged
     solution; the precipitated surface-modified zinc oxide was
     separated, dried and disintegrated; 784 g product was obtained. A sunscreen contained (weight/weight%): Cetiol OE 10; Cetiol S 10; Lanette O 4.5; Eumulgin
     B2 2; Monomuls 60-35C 2; Basysilon M 350 0.5; Phenonip 1; zinc
     oxide (surface modified with isostearic acid) 8; zinc
     oxide (surface modified with polyglycol diacid) 8; water to 100.
     zinc oxide nanoparticle surface modification
     isostearate redispersion cosmetics conductor
     Cosmetics
        (foams; redispersible nanoparticulate zinc
        oxide powder with surface coating for use in cosmetics and for
        elec. conducting surfaces)
     Cosmetics
        (gels; redispersible nanoparticulate zinc
        oxide powder with surface coating for use in cosmetics and for
        elec. conducting surfaces)
     Cosmetics
        (lotions; redispersible nanoparticulate zinc
        oxide powder with surface coating for use in cosmetics and for
        elec. conducting surfaces)
    Cosmetics
        (moisturizers; redispersible nanoparticulate zinc
        oxide powder with surface coating for use in cosmetics and for
        elec. conducting surfaces)
     Solvents
        (organic; redispersible nanoparticulate zinc
        oxide powder with surface coating for use in cosmetics and for
        elec. conducting surfaces)
    Cosmetics
        (powders; redispersible nanoparticulate zinc
        oxide powder with surface coating for use in cosmetics and for
        elec. conducting surfaces)
    Antimicrobial agents
    Antioxidants
    Deodorants
    Dispersion (of materials)
    Electric conductors
    Milling (size reduction)
    Nanoparticles
    Pigments, nonbiological
    Polarity
    Precipitation (chemical)
    Stability
    Sunscreens
    Surface treatment
    Temperature
        (redispersible nanoparticulate zinc oxide
        powder with surface coating for use in cosmetics and for elec.
        conducting surfaces)
    Cosmetics
        (sticks; redispersible nanoparticulate zinc
        oxide powder with surface coating for use in cosmetics and for
        elec. conducting surfaces)
    1406-18-4, Vitamine
    RL: COS (Cosmetic use); BIOL (Biological study); USES (Uses)
        (redispersible nanoparticulate zinc oxide
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powder with surface coating for use in cosmetics and for elec.
         conducting surfaces)
      1314-13-2, Zinc oxide, biological studies
TΤ
      26009-03-0D, Polyglycolic acid, derivs. 26124-68-5D, Polyglycolic acid,
                  27306-90-7 30399-84-9, Isostearic acid 53563-70-5
      RL: COS (Cosmetic use); TEM (Technical or engineered material use); BIOL
      (Biological study); USES (Uses)
          (redispersible nanoparticulate zinc oxide
         powder with surface coating for use in cosmetics and for elec.
         conducting surfaces)
IT
      56-23-5, Tetrachloromethane, uses 60-29-7, Diethylether, uses 67-66-3,
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      76-16-4, Hexafluoroethane 78-87-5, 1,2-Dichloropropane 79-34-5,
      1,1,2,2-Tetrachloroethane 95-47-6, o-Xylene, uses 98-06-6, tert-Butyl
      benzene 98-82-8, Isopropylbenzene 100-41-4, Ethylbenzene, uses 101-84-8, Diphenylether 105-58-8, Diethyl carbonate 106-42-3,
      p-Xylene, uses 107-04-0, 1-Bromo-2-chloroethane 107-06-2,
     1,2-Dichloroethane, uses 108-20-3, Diisopropylether 108-38-3, m-Xylene, uses 108-88-3, Toluene, uses 109-66-0, Pentane, uses
      109-99-9, Tetrahydrofuran, uses 110-54-3, Hexane, uses 111-65-9,
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      Pentafluoroethane 540-67-0, Methylethylether 616-38-6, Dimethyl
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      RL: TEM (Technical or engineered material use); USES (Uses)
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         conducting surfaces)
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                              2004:509968 CAPLUS
DOCUMENT NUMBER:
                              141:59224
TITLE:
                             Redispersible nanoparticulate zinc
                              oxide powder with surface coating for use in
                              cosmetics and for electrical conducting surfaces
INVENTOR(S):
                             Elsaesser, Ralf; Kliss, Rainer; Hahn, Horst; Kropf,
                              Christian; Berber, Mete; Carulla, Victor Bulto
                             Sustech GmbH & Co. KG, Germany
PATENT ASSIGNEE(S):
SOURCE:
                              Ger. Offen., 19 pp.
                             CODEN: GWXXBX
DOCUMENT TYPE:
                              Patent
LANGUAGE:
                             German
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
      PATENT NO.
                             KIND
                                     DATE
                                             APPLICATION NO.
                                                                               DATE
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     DE 10257388
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     WO 2004052327 A2
WO 2004052327 A3
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                                      20040624
                                                                               20031126
                                     20041014

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    RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR

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                                                   AU 2003-288172
                                                                               20031126
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                                                                               20031126
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DE 2002-10257388 A 20021206 WO 2003-EP13286 W 20031126
PRIORITY APPLN. INFO.:
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OTHER SOURCE(S): MARPAT 141:59224

AB The invention concerns nanoparticular zinc oxide that is surface-modified with an organic acid, e.g. isostearic acid or

polyglycolic acid derivs. The surface-treated zinc oxide can be redispersed in polar and apolar organic solvents. Particles for elec. conducting surfaces and cosmetics are prepared Thus zinc oxide was precipitated by dissolving 1110 g zinc chloride in 11.1 L methanol and adding 651 g sodium hydroxide pellets under stirring and maintaining room temperature The NaCl/ZnO mixture was separated, followed by washing and resuspension in 11 L THF. 385 G isostearic acid were added; after 3 h stirring was stopped; sodium chloride settled and coated zinc oxide was contained in the clear supernatant. The clear supernatant was separated, the solvent evaporated and

the

raw product reconstructed in toluene; the residual sodium chloride was removed by centrifugation. 4L ethanol (96%) were added to the centrifuged solution; the precipitated surface-modified zinc oxide was separated, dried and disintegrated; 784 g product was obtained. A sunscreen contained (weight/weight%): Cetiol OE 10; Cetiol S 10; Lanette O 4.5; Eumulgin B2 2; Monomuls 60-35C 2; Basysilon M 350 0.5; Phenonip 1; zinc oxide (surface modified with isostearic acid) 8; zinc oxide (surface modified with polyglycol diacid) 8; water to 100.

- L5 ANSWER 9 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
- AN 2003:765357 CAPLUS
- DN 139:389898
- ED Entered STN: 01 Oct 2003
- TI Magnetic Quantum Dots: Synthesis, Spectroscopy, and Magnetism of Co2+- and Ni2+-Doped ZnO Nanocrystals
- AU Schwartz, Dana A.; Norberg, Nick S.; Nguyen, Quyen P.; Parker, Jason M.; Gamelin, Daniel R.
- CS Department of Chemistry, University of Washington, Seattle, WA, 98195-1700, USA
- SO Journal of the American Chemical Society (2003), 125(43), 13205-13218 CODEN: JACSAT; ISSN: 0002-7863
- PB American Chemical Society
- DT Journal
- LA English
- CC 77-1 (Magnetic Phenomena)
 Section cross-reference(s): 66
- AB The authors report a method for the preparation of colloidal ZnO
 -diluted magnetic semiconductor quantum dots (DMS-QDs) by alkaline-activated hydrolysis and condensation of Zn acetate solns. in DMSO. Mechanistic studies reveal that Co2+ and Ni2+ dopants inhibit nucleation and growth of ZnO nanocrystals. In particular, dopants are quant. excluded from the critical nuclei but are incorporated nearly isotropically during subsequent growth of the nanocrystals. The smaller nanocrystal diams. that result upon doping are explained by the Gibbs-Thompson relation between lattice strain and crystal solubility. The authors describe methods for cleaning the nanocrystal surfaces of exposed dopants and for redispersion of the final DMS-QDs. Homogeneous substitutional doping is verified by high-resolution low-temperature electronic absorption and MCD

spectroscopies. A giant Zeeman effect is observed in the band gap transition of Co2+:ZnO DMS-QDs. MCD and Zeeman spectroscopies were used to quantify the magnitude of the p-d exchange interaction (NO β) that gives rise to this effect. NO β values of -2.3 \pm 0.3 eV (-18,500 cm-1) for Co2+:ZnO and -4.5 \pm 0.6 eV (-36,300 cm-1) for Ni2+: ZnO were determined Ligand-to-metal charge-transfer transitions are observed in the MCD spectra of both Co2+:ZnO and Ni2+:ZnO DMS-QDs and are analyzed in the context of an optical electronegativity model. The importance of these charge-transfer states in determining NO β is discussed. Ferromagnetism with TC > 350 K is observed in aggregated nanocrystals of Co2+:ZnO that unambiguously demonstrates the existence of intrinsic high-TC ferromagnetism in this class of DMSs.

- ST quantum dot spectroscopy magnetism cobalt nickel zinc oxide nanocrystal
- IT Zeeman effect

```
(giant Zeeman effect observed in band gap transition of Co2+:ZnO
        nanocryst. magnetic semiconductor quantum dots)
TΤ
     Exchange interaction
     Magnetooptical effect
        (in Co2+- and Ni2+-doped ZnO nanocryst. magnetic quantum dots
        in DMSO)
ΙT
     Ferromagnetism
        (in Co2+:ZnO aggregated nanocryst. magnetic semiconductor
        quantum dots)
ΙT
     Electron transfer
        (ligand-to-metal charge-transfer transitions observed in the MCD spectra
        of both Co2+: ZnO and Ni2+: ZnO magnetic
        semiconductor quantum dots)
IT
     Colloids
     Dopina
     Magnetic semiconductor materials
     Nanocrystals
        (magnetic quantum dots: synthesis, spectroscopy, and magnetism of Co2+-
        and Ni2+-doped ZnO nanocryst. magnetic quantum dots in DMSO)
TΤ
     Magnetic circular dichroism
        (of ZnO nanocryst. magnetic semiconductor quantum dots in
        conformation of cobalt and nickel doping)
IΤ
     Band gap
        (transition; giant Zeeman effect observed in band gap transition of Co2+:
        ZnO nanocryst. magnetic semiconductor quantum dots)
IT
     14701-22-5, Nickel(2+), properties
                                          22541-53-3, Cobalt(2+), properties
     RL: MOA (Modifier or additive use); PRP (Properties); USES (Uses)
        (magnetic quantum dots: synthesis, spectroscopy, and magnetism of Co2+-
        and Ni2+-doped ZnO nanocryst. magnetic quantum dots in DMSO)
     1314-13-2, Zinc oxide (ZnO), properties
ΙT
     RL: PEP (Physical, engineering or chemical process); PRP (Properties); PYP
     (Physical process); PROC (Process)
        (magnetic quantum dots: synthesis, spectroscopy, and magnetism of Co2+-
        and Ni2+-doped ZnO nanocryst. magnetic quantum dots in DMSO)
RE.CNT
              THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD
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ACCESSION NUMBER:
                             2003:765357 CAPLUS
DOCUMENT NUMBER:
                             139:389898
TITLE:
                             Magnetic Quantum Dots: Synthesis, Spectroscopy, and
                             Magnetism of Co2+- and Ni2+-Doped ZnO
                             Nanocrystals
AUTHOR(S):
                             Schwartz, Dana A.; Norberg, Nick S.; Nguyen, Quyen P.;
                             Parker, Jason M.; Gamelin, Daniel R.
CORPORATE SOURCE:
                             Department of Chemistry, University of Washington,
                             Seattle, WA, 98195-1700, USA
SOURCE:
                             Journal of the American Chemical Society (2003),
                             125(43), 13205-13218
                             CODEN: JACSAT; ISSN: 0002-7863
PUBLISHER:
                             American Chemical Society
DOCUMENT TYPE:
                             Journal
LANGUAGE:
                             English
     The authors report a method for the preparation of colloidal ZnO
     -diluted magnetic semiconductor quantum dots (DMS-QDs) by alkaline-activated
     hydrolysis and condensation of Zn acetate solns. in DMSO. Mechanistic
     studies reveal that Co2+ and Ni2+ dopants inhibit nucleation and growth of
     ZnO nanocrystals. In particular, dopants are quant. excluded from
     the critical nuclei but are incorporated nearly isotropically during
     subsequent growth of the nanocrystals. The smaller nanocrystal diams.
     that result upon doping are explained by the Gibbs-Thompson relation
     between lattice strain and crystal solubility The authors describe methods for
     cleaning the nanocrystal surfaces of exposed dopants and for
     redispersion of the final DMS-QDs. Homogeneous substitutional
     doping is verified by high-resolution low-temperature electronic absorption
and MCD
     spectroscopies. A giant Zeeman effect is observed in the band gap transition
     of Co2+:ZnO DMS-QDs. MCD and Zeeman spectroscopies were used to
     quantify the magnitude of the p-d exchange interaction (N0\beta) that
     gives rise to this effect. NO\beta values of -2.3 \pm 0.3 eV (-18,500
     cm-1) for Co2+:ZnO and -4.5 \pm 0.6 eV (-36,300 cm-1) for Ni2+:
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ZnO were determined Ligand-to-metal charge-transfer transitions are observed in the MCD spectra of both Co2+:ZnO and Ni2+:ZnO DMS-QDs and are analyzed in the context of an optical electronegativity model. The importance of these charge-transfer states in determining NO β is discussed. Ferromagnetism with TC > 350 K is observed in aggregated nanocrystals of Co2+:ZnO that unambiguously demonstrates the existence of intrinsic high-TC ferromagnetism in this class of DMSs.

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L_5
     ANSWER 10 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
ΑN
     2003:656693 CAPLUS
DN
     139:182379
ED
     Entered STN: 22 Aug 2003
     Synthesis and use of nanoscale rutile or nanoscale metal oxides as
TI
     photo-inactive transparent fillers
ΙN
     Nonninger, Ralph
PA
     ITN-Nanovation G.m.b.H., Germany
     PCT Int. Appl., 26 pp.
SO
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LA
    German
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    ICM C01G023-053
     ICS C01G025-02; C01G023-00; C01G001-02
CC
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CLASS
PATENT NO.
               CLASS PATENT FAMILY CLASSIFICATION CODES
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WO 2003068682
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                ICS
                       C01G025-02; C01G023-00; C01G001-02
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                  FTERM
                         4G047/CA02; 4G047/CA05; 4G047/CB06; 4G047/CB09;
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 US 2005233146
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                         [I,C*]; C01B0013-36 [I,A]; C01G0001-02 [I,C*];
                         C01G0001-02 [I,A]; C01G0023-00 [I,C*]; C01G0023-00
                         [I,A]; C01G0023-053 [I,A]; C01G0025-00 [I,C*];
                         C01G0025-02 [I,A]
                 NCL
                         428/402.000; 423/598.000; 502/350.000
AB
     Nanoscale rutile (TiO2) is prepared from amorphous TiO2 by mixing an alc.
     solution with a titanium alcoholate (and, optionally, a second alcoholate)
    and adding water and acid. The amorphous TiO2 is then either recovered by
     removal of solvent or by crystallization from the solvent, and then
redispersed in
     water in the presence of a tin salt. A thermal or hydrothermal
     post-treatment yields rutile or an oxide that can be redispersed to the
     desired particle size (5-20 nm). The redispersed rutile or the obtained
     oxide, with primary particle size 5-20 nm, can be incorporated into all
     organic matrixes so that they remain transparent. Photocatalytic activity is suppressed by lattice doping with trivalent ions. A variation of the
     alcoholates allows production of the following components: ZrO2, ZnO
     , SnO2, ATO (antimony tin oxide), In2O3, ITO (indium tin oxide), Y2O3,
     BaO, CaO, MgO, CeO2, and BaTiO3.
     nanoparticle rutile photoinactive filler; metal oxide nanoparticle
ST
     synthesis filler
ΙT
     Titanates
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (alkoxides, titanium source; synthesis and use of nanoscale rutile or
        nanoscale metal oxides as photo-inactive transparent fillers)
ΙT
     Zirconates
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
       (alkoxides, zirconium source; synthesis and use of nanoscale rutile or
        nanoscale metal oxides as photo-inactive transparent fillers)
ΙT
     Metal alkoxides
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (aluminum, aluminum source; synthesis and use of nanoscale rutile or
        nanoscale metal oxides as photo-inactive transparent fillers)
ΙT
     Metal alkoxides
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
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```
(metal source; synthesis and use of nanoscale rutile or nanoscale metal
        oxides as photo-inactive transparent fillers)
     Alcohols, uses
TT
     RL: NUU (Other use, unclassified); USES (Uses)
        (solvents; synthesis and use of nanoscale rutile or nanoscale metal
        oxides as photo-inactive transparent fillers)
ΙT
     Nanoparticles
        (synthesis and use of nanoscale rutile or nanoscale metal oxides as
        photo-inactive transparent fillers)
ΙT
     Metal alkoxides
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (titanium, titanium source; synthesis and use of nanoscale rutile or
        nanoscale metal oxides as photo-inactive transparent fillers)
ΙT
     Fillers
        (transparent; synthesis and use of nanoscale rutile or nanoscale metal
        oxides as photo-inactive transparent fillers)
IT
     Metal alkoxides
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (yttrium; synthesis and use of nanoscale rutile or nanoscale metal
        oxides as photo-inactive transparent fillers)
TT
     Metal alkoxides
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (zirconium, zirconium source; synthesis and use of nanoscale rutile or
        nanoscale metal oxides as photo-inactive transparent fillers)
IT
     7429-90-5D, Aluminum, alcoholates
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (aluminum source; synthesis and use of nanoscale rutile or nanoscale
        metal oxides as photo-inactive transparent fillers)
     1304-28-5P, Barium oxide, preparation 1305-78-8P, Ca preparation 1306-38-3P, Cerium dioxide, preparation
IT
                                              1305-78-8P, Calcium oxide,
                                                               1309-48-4P,
     Magnesium oxide, preparation
                                     1312-43-2P, Indium oxide (In2O3)
     1314-13-2P, Zinc oxide, preparation
                                            1314-36-9P,
     Yttrium oxide (Y2O3), preparation
                                         1317-80-2P, Rutile
                                                                12047-27-7P,
     Barium titanate (BatiO3), preparation 12673-86-8P, Antimony tin oxide 18282-10-5P, Tin dioxide 50926-11-9P, Indium tin oxide
     RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP
     (Preparation)
        (nanoscale; synthesis and use of nanoscale rutile or nanoscale metal
        oxides as photo-inactive transparent fillers)
TΤ
     75-09-2, Dichloromethane, uses
                                      109-99-9, Tetrahydrofuran, uses
     1344-13-4, Tin chloride
                                7440-31-5D, Tin, salts 7647-14-5, Sodium
     chloride, uses
                       10031-62-6, Tin sulfate 12125-02-9, Ammonium chloride,
            14536-20-0
     RL: NUU (Other use, unclassified); USES (Uses)
        (redispersion solns. containing; synthesis and use of nanoscale
        rutile or nanoscale metal oxides as photo-inactive transparent fillers)
IT
     7440-32-6D, Titanium, alcoholates
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (titanium source; synthesis and use of nanoscale rutile or nanoscale
        metal oxides as photo-inactive transparent fillers)
IT
     7440-65-5D, Yttrium, alcoholates
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (yttrium source; synthesis and use of nanoscale rutile or nanoscale
        metal oxides as photo-inactive transparent fillers)
IT
     7440-67-7D, Zirconium, alcoholates
     RL: CPS (Chemical process); PEP (Physical, engineering or chemical
     process); PROC (Process)
        (zirconium source; synthesis and use of nanoscale rutile or nanoscale
        metal oxides as photo-inactive transparent fillers)
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RE.CNT 5
               THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE
(1) Arney, D; WO 0006495 A 2000 CAPLUS
(2) Chemseddine, A; EUR J INORG CHEM 1999, P235 CAPLUS
(3) Cheng, H; CHEM MATER 1995, V7(4), P663 CAPLUS (4) Kronos Int Inc; DE 4105345 A 1992 CAPLUS
(5) Lee, S; J PHOCHEM AND PHOTOBIOL A: CHEMISTRY 2001, V146, P121 CAPLUS
ACCESSION NUMBER:
                          2003:656693 CAPLUS
DOCUMENT NUMBER:
                          139:182379
TITLE:
                           Synthesis and use of nanoscale rutile or nanoscale
                           metal oxides as photo-inactive transparent fillers
INVENTOR(S):
                           Nonninger, Ralph
                         ITN-Nanovation G.m.b.H., Germany
PATENT ASSIGNEE(S):
SOURCE:
                           PCT Int. Appl., 26 pp.
                           CODEN: PIXXD2
DOCUMENT TYPE:
                           Patent
LANGUAGE:
                           German
FAMILY ACC. NUM. COUNT: 1
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     WO 2003068682 A1 20030821 WO 2002-DE3188 20020830
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A1 20041117 EP 2002-806790 20020830
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      JP 2005528309
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      JP 2003-567825

      US 2005233146
      A1
      20051020
      US 2005-503627

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                                               DE 2002-10205920 A 20020212
WO 2002-DE3188 W 20020830
PRIORITY APPLN. INFO.:
     Nanoscale rutile (TiO2) is prepared from amorphous TiO2 by mixing an alc.
AB
     solution with a titanium alcoholate (and, optionally, a second alcoholate)
     and adding water and acid. The amorphous TiO2 is then either recovered by
     removal of solvent or by crystallization from the solvent, and then
redispersed in
     water in the presence of a tin salt. A thermal or hydrothermal
     post-treatment yields rutile or an oxide that can be redispersed to the
     desired particle size (5-20 nm). The redispersed rutile or the obtained
     oxide, with primary particle size 5-20 nm, can be incorporated into all
     organic matrixes so that they remain transparent. Photocatalytic activity is
     suppressed by lattice doping with trivalent ions. A variation of the
     alcoholates allows production of the following components: ZrO2, ZnO
     , SnO2, ATO (antimony tin oxide), In2O3, ITO (indium tin oxide), Y2O3,
     BaO, CaO, MgO, CeO2, and BaTiO3.
L5
     ANSWER 11 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
     2000:593025 CAPLUS
ΑN
DN
     133:179813
     Entered STN: 25 Aug 2000
ED
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Preparation of precipitated zinc oxide nanoparticles

Womelsdorf, Hermann; Hoheisel, Werner; Passing, Gerd

by alkaline hydrolysis, flocculation, gel formation, and sol

ΤI

ΙN

redispersion

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PA
     Bayer A.-G., Germany
SO
     Ger. Offen., 4 pp.
     CODEN: GWXXBX
DT
     Patent
LA
     German
IC
     ICM C01G009-02
     ICS C08K003-22; A61K007-40
CC
     49-3 (Industrial Inorganic Chemicals)
     Section cross-reference(s): 38
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     DE 19907704 A1 20000824 DE 1999-19907704 19990223
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                     T 20040915 AT 2000-907548
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B 20041221 TW 2000-89102918
     AT 275166
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US 6710091 B1 20040323
PRAI DE 1999-19907704 A 19990223
WO 2000-EP1116 W 20000211
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CLASS
                  CLASS PATENT FAMILY CLASSIFICATION CODES
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                          C01G0009-02 [ICM,7]; C01G0009-00 [ICM,7,C*];
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                         [I,A]; C08K0003-00 [I,C*]; C08K0003-22 [I,A]
                 NCL
                         516/033.000; 423/101.000; 423/102.000; 423/104.000;
                         423/622.000; 516/088.000; 516/098.000
                         A61K008/04H; A61K008/27; A61Q017/04; C01G009/02;
                 ECLA
                         C08K003/22
     Precipitated zinc oxide nanoparticles, with average particle
AΒ
     diameter <10 nm, are easily redispersible, primarily as a sol, in
     organic solvents, water, solvent mixts., and solvent mixts. containing
     surface-modifying substances. The ZnO nanoparticles are prepared
     by alkaline hydrolysis in alc. solution, in which the ppts. formed during
     hydrolysis are aged until the ZnO is completely flocculated,
     which is then compressed to a gel form, and separated from the supernatant
     that contains the byproducts. The ZnO sols are prepared from the
     gel-like precipitated zinc oxide by treatment of the ppts.
     with an organic solvent, a solvent mixture, or water, or a water-alc. mixture
     containing a surface-modifying substance. Suitable solvents are methylene
     chloride or chloroform. These precipitated ZnO nanoparticles are used
     for matrix modification of polymers, resins, and coatings as well as for
     the UV protection of polymers, resins, coatings, and UV-sensitive organic
     pigments.
     zinc oxide nanoparticle prepn use; sol pptn
ST
     redispersion zinc oxide nanoparticle
IT
     Hydrolysis
        (base, of zinc salts; preparation of precipitated zinc oxide
        nanoparticles by alkaline hydrolysis, flocculation, gel formation, and sol
        redispersion)
IT
     Precipitation (chemical)
        (of aged zinc oxide flocculates; preparation of precipitated
        zinc oxide nanoparticles by alkaline hydrolysis,
        flocculation, gel formation, and sol redispersion)
ΙT
     Pigments, nonbiological
        (organic, zinc oxide nanoparticles as UV stabilizers
        for; preparation of precipitated zinc oxide nanoparticles by
        alkaline hydrolysis, flocculation, gel formation, and sol
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(preparation of precipitated zinc oxide nanoparticles by alkaline hydrolysis, flocculation, gel formation, and sol redispersion

redispersion)

Flocculation

IT

```
IT
     Polymers, miscellaneous
     RL: MSC (Miscellaneous)
        (zinc oxide nanoparticles as matrix modifiers for;
        preparation of precipitated zinc oxide nanoparticles by alkaline
        hydrolysis, flocculation, gel formation, and sol redispersion
TT
     Coating materials
     Fillers
     UV stabilizers
        (zinc oxide nanoparticles; preparation of precipitated
        zinc oxide nanoparticles by alkaline hydrolysis,
        flocculation, gel formation, and sol redispersion)
IT
     Colloids
     Nanoparticles
     Sols
        (zinc oxide; preparation of precipitated zinc
        oxide nanoparticles by alkaline hydrolysis, flocculation, gel
        formation, and sol redispersion)
     5970-45-6, Zinc acetate dihydrate
ΙT
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (alkaline hydrolysis of; preparation of precipitated zinc oxide
        nanoparticles by alkaline hydrolysis, flocculation, gel formation, and sol
        redispersion)
IT
     1310-58-3, Potassium hydroxide, uses
     RL: NUU (Other use, unclassified); USES (Uses)
        (alkaline hydrolysis with; preparation of precipitated zinc oxide
        nanoparticles by alkaline hydrolysis, flocculation, gel formation, and sol
        redispersion)
ΙT
     1314-13-2P, Zinc oxide, preparation
     RL: NUU (Other use, unclassified); PRP (Properties); SPN (Synthetic
     preparation); PREP (Preparation); USES (Uses)
        (preparation of precipitated zinc oxide nanoparticles by alkaline
        hydrolysis, flocculation, gel formation, and sol redispersion
ΙT
     67-56-1, Methanol, uses
     RL: NUU (Other use, unclassified); USES (Uses)
        (solvent, alkaline hydrolysis in presence of; preparation of precipitated
zinc
        oxide nanoparticles by alkaline hydrolysis, flocculation, gel
        formation, and sol redispersion)
     67-66-3, Chloroform, uses
                                75-09-2, Methylene chloride, uses
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        (solvent, for preparation of zinc oxide sol; preparation of
        precipitated zinc oxide nanoparticles by alkaline hydrolysis,
        flocculation, gel formation, and sol redispersion)
ACCESSION NUMBER:
                         2000:593025 CAPLUS
DOCUMENT NUMBER:
                         133:179813
TITLE:
                         Preparation of precipitated zinc
                         oxide nanoparticles by alkaline hydrolysis,
                         flocculation, gel formation, and sol
                         redispersion
INVENTOR(S):
                         Womelsdorf, Hermann; Hoheisel, Werner; Passing, Gerd
PATENT ASSIGNEE(S):
                         Bayer A.-G., Germany
                         Ger. Offen., 4 pp.
SOURCE:
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DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         German
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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WO 2000-EP1116

WO 2000050503

A1

20000831

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             SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM,
             AZ, BY, KG, KZ, MD, RU, TJ, TM
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PRIORITY APPLN. INFO.:
                                            DE 1999-19907704
                                                                A 19990223
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     Precipitated zinc oxide nanoparticles, with average particle
AB
     diameter <10 nm, are easily redispersible, primarily as a sol, in
     organic solvents, water, solvent mixts., and solvent mixts. containing
     surface-modifying substances. The ZnO nanoparticles are prepared
     by alkaline hydrolysis in alc. solution, in which the ppts. formed during
alkaline
     hydrolysis are aged until the ZnO is completely flocculated,
     which is then compressed to a gel form, and separated from the supernatant
     that contains the byproducts. The ZnO sols are prepared from the
     gel-like precipitated zinc oxide by treatment of the ppts.
     with an organic solvent, a solvent mixture, or water, or a water-alc. mixture
     containing a surface-modifying substance. Suitable solvents are methylene
     chloride or chloroform. These precipitated ZnO nanoparticles are used
     for matrix modification of polymers, resins, and coatings as well as for
     the UV protection of polymers, resins, coatings, and UV-sensitive organic
     pigments.
L5
     ANSWER 12 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
ΑN
     1998:406685 CAPLUS
DΝ
     129:141102
ED
     Entered STN: 03 Jul 1998
     Synthesis and Growth of ZnO Nanoparticles
TI
ΑU
     Meulenkamp, Eric A.
CS
     Philips Research Laboratories, Eindhoven, 5656 AA, Neth.
SO
     Journal of Physical Chemistry B (1998), 102(29), 5566-5572
     CODEN: JPCBFK; ISSN: 1089-5647
PΒ
     American Chemical Society
DT
     Journal
LA
     English
CC
     66-4 (Surface Chemistry and Colloids)
AΒ
     ZnO nanoparticles in the size range from 2 to 7 nm were prepared
     by addition of LiOH to an EtOH-Zn acetate solution This method [Spanhel, L.;
     Anderson, M. A. J. Am. Chemical Society 1991, 113, 2826] was modified and
     extended at several points. The synthesis of very small ZnO
     nuclei was simplified. Aging of particles was governed by temperature, the H2O
     content, and the presence of reaction products. H2O and acetate induced
     considerably accelerated particle growth. Growth could almost be stopped
     by removal of these species (washing). Washing consisted of repeated
     precipitation of ZnO by addition of alkanes such as heptane, removal of the
     supernatant, and redispersion in EtOH. The aging
     characteristics are interpreted in terms of the concentration of dissolved ZnII
     species and reactions known in sol-gel chemical These findings present a
     better-defined and more versatile procedure for production of clean
     ZnO sols of readily adjustable particle size. Such sols are of
     particular interest for studies of elec. and optical properties of
```

W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU,

```
ZnO nanoparticle films. For example, films exhibiting >99%
     transparency in the visible region could only be obtained by deposition
     from a washed soluble
ST
     sol gel processing zinc oxide nanoparticle
ΙT
     Particle size
        (of zinc oxide nanoparticle in sol)
     UV absorption
IT
        (of zinc oxide nanoparticle sols)
TT
     Band gap
        (optical; of zinc oxide nanoparticle sols)
IT
     Dispersion (of materials)
        (re-; precipitation and redispersion of zinc oxide
        nanoparticles)
ΙT
     Nanoparticles
     Sol-gel processing
        (synthesis and growth of ZnO nanoparticles by sol gel
        processing)
     1314-13-2, Zinc oxide (ZnO), properties
IT
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (synthesis and growth of ZnO nanoparticles by sol gel
        processing)
RE.CNT
        32
              THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS RECORD
RE
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ACCESSION NUMBER:
                         1998:406685 CAPLUS
DOCUMENT NUMBER:
                         129:141102
TITLE:
                         Synthesis and Growth of ZnO Nanoparticles
                         Meulenkamp, Eric A.
AUTHOR(S):
CORPORATE SOURCE:
                         Philips Research Laboratories, Eindhoven, 5656 AA,
SOURCE:
                         Journal of Physical Chemistry B (1998), 102(29),
                         5566-5572
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CODEN: JPCBFK; ISSN: 1089-5647

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

ZnO nanoparticles in the size range from 2 to 7 nm were prepared by addition of LiOH to an EtOH-Zn acetate solution This method [Spanhel, L.; Anderson, M. A. J. Am. Chemical Society 1991, 113, 2826] was modified and extended at several points. The synthesis of very small ZnO nuclei was simplified. Aging of particles was governed by temperature, the H2O content, and the presence of reaction products. H2O and acetate induced considerably accelerated particle growth. Growth could almost be stopped by removal of these species (washing). Washing consisted of repeated precipitation of ZnO by addition of alkanes such as heptane, removal of the supernatant, and redispersion in EtOH. The aging characteristics are interpreted in terms of the concentration of dissolved ZnII species and reactions known in sol-gel chemical These findings present a better-defined and more versatile procedure for production of clean ZnO sols of readily adjustable particle size. Such sols are of particular interest for studies of elec. and optical properties of ZnO nanoparticle films. For example, films exhibiting >99% transparency in the visible region could only be obtained by deposition from a washed soluble

- L5 ANSWER 13 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
- ΑN 1998:374162 CAPLUS
- DN 129:9090
- ED Entered STN: 19 Jun 1998
- ΤI Effects of reduction and regeneration conditions on the activity of CuO-ZnO catalysts
- ΑU Quincoces, C. E.; Amadeo, N. E.; Gonzalez, M. G.
- CS UNLP-CONICET, La Plata, 1900, Argent.
- SO Studies in Surface Science and Catalysis (1997), 111(Catalyst Deactivation 1997), 535-541 CODEN: SSCTDM; ISSN: 0167-2991
 - Elsevier Science B.V.
- PΒ
- DTJournal
- LA English
- CC 67-1 (Catalysis, Reaction Kinetics, and Inorganic Reaction Mechanisms)
- The influence of thermal and hydrothermal treatments and reduction processes AΒ on the activity of CuO-ZnO catalysts was studied.

Temperature-programmed reduction (TPR) was used in the study of catalyst activation

with different reduction mixts. A mixture of H2 and N2 in the range of 500 to 573 K was the most effective reducing agent. High temperature, water vapor and a high partial pressure of hydrogen enhance Cu sintering and Cu-Zn alloy formation with a decrease in catalytic activity. XRD anal. of reactivated catalysts showed that their deactivation can be related to the formation of surface spinel species. Reoxidn. at 623 K and subsequent reduction of the aged catalyst led to metal redispersion but not catalyst regeneration.

- redn activity copper zinc oxide catalyst; regeneration ST activity copper zinc oxide catalyst
- ΙT

Water gas shift reaction catalysts

(effects of reduction and regeneration conditions on activity of CuO-ZnO catalysts)

ITReduction

> (thermal; effects of reduction and regeneration conditions on activity of CuO-ZnO catalysts)

ΙT 11143-56-9

> RL: CAT (Catalyst use); FMU (Formation, unclassified); PEP (Physical, engineering or chemical process); FORM (Formation, nonpreparative); PROC (Process); USES (Uses)

(effects of reduction and regeneration conditions on activity of CuO-ZnO catalysts)

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ΤТ
     1314-13-2, Zinc oxide, uses
                                    1317-38-0, Copper
                         1344-28-1, Alumina, uses
     oxide(cuo), uses
     RL: CAT (Catalyst use); PEP (Physical, engineering or chemical process);
     PRP (Properties); PROC (Process); USES (Uses)
         (effects of reduction and regeneration conditions on activity of CuO-
        ZnO catalysts)
     7727-37-9, Nitrogen, processes
IT
     RL: PEP (Physical, engineering or chemical process); PROC (Process)
         (effects of reduction and regeneration conditions on activity of CuO-
        ZnO catalysts)
     1333-74-0, Hydrogen, reactions
ΙT
     RL: RCT (Reactant); RACT (Reactant or reagent)
         (effects of reduction and regeneration conditions on activity of CuO-
        ZnO catalysts)
RE.CNT
               THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD
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(14) Young, P; Chem Eng Prog V69
ACCESSION NUMBER:
                          1998:374162 CAPLUS
DOCUMENT NUMBER:
                          129:9090
TITLE:
                          Effects of reduction and regeneration conditions on
                          the activity of CuO-ZnO catalysts
                          Quincoces, C. E.; Amadeo, N. E.; Gonzalez, M. G.
AUTHOR(S):
                          UNLP-CONICET, La Plata, 1900, Argent.
CORPORATE SOURCE:
SOURCE:
                          Studies in Surface Science and Catalysis (1997),
                          111(Catalyst Deactivation 1997), 535-541
                          CODEN: SSCTDM; ISSN: 0167-2991
PUBLISHER:
                          Elsevier Science B.V.
DOCUMENT TYPE:
                          Journal
LANGUAGE:
                          English
     The influence of thermal and hydrothermal treatments and reduction processes
     on the activity of CuO-ZnO catalysts was studied.
     Temperature-programmed reduction (TPR) was used in the study of catalyst
activation
     with different reduction mixts. A mixture of H2 and N2 in the range of 500 to
     573 K was the most effective reducing agent. High temperature, water vapor and
     a high partial pressure of hydrogen enhance Cu sintering and Cu-Zn alloy
     formation with a decrease in catalytic activity. XRD anal. of reactivated
     catalysts showed that their deactivation can be related to the formation
     of surface spinel species. Reoxidn. at 623 K and subsequent reduction of the
     aged catalyst led to metal redispersion but not catalyst
     regeneration.
L5
     ANSWER 14 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
ΑN
     1995:742638 CAPLUS
DN
     123:115526
ED
     Entered STN: 18 Aug 1995
     Blue pigment slurries and their manufacture
TI
ΙN
     Mizukami, Tomohito; Funayama, Chihito; Terada, Hiromi; Nishio, Akira
PA
     Kasei Optonix, Japan; Dainichiseika Color Chem
     Jpn. Kokai Tokkyo Koho, 2 pp.
SO
     CODEN: JKXXAF
DT
     Patent
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LA
     Japanese
IC
     ICM C09C001-30
     ICS C09C001-30; C09K011-02
     42-6 (Coatings, Inks, and Related Products)
CC
FAN.CNT 1
                               DATE APPLICATION NO. DATE
     PATENT NO.
                       KIND
                               DATE
                         ____
                                19950314 JP 1993-148931
PI JP 07070466
PRAI JP 1993-148931
                         Α
                                                                  19930621
                                19930621
CLASS
 PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES
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                ICM
ICS
 JP 07070466
                       C09C001-30
                       C09C001-30; C09K011-02
                 IPCI C09C0001-30 [ICM, 6]; C09C0001-30 [ICS, 6]; C09C0001-28
                       [ICS, 6, C*]; C09K0011-02 [ICS, 6]
                        C09C0001-30 [I,A]; C09C0001-28 [I,C*]; C09C0001-28
                 IPCR
                        [I,A]; C09K0011-02 [I,C*]; C09K0011-02 [I,A]
AΒ
     Title slurries with good re-dispersibility are prepared by wet dispersing
     CoO.ZnO.SiO2 particles in the presence of organic acids or H3PO4.
     Ball-milling 1.2-\mu m CoO .ZnO.SiO2 particles with water and
     citric acid at 110 rpm for 5 h to form 0.4-µm particles and setting at
     room temperature for 24 h gave soft and re-dispersible precipitate
     redispersibility cobalt zinc silicon oxide pigment; org acid manuf
     redispersible blue pigment; citric acid manuf
     redispersible blue pigment
ΙT
     Pigments
        (manufacture of re-dispersible blue cobalt zinc silicon oxide pigments)
ΙT
     Carboxylic acids, uses
     RL: NUU (Other use, unclassified); USES (Uses)
        (manufacture of re-dispersible blue cobalt zinc silicon oxide pigments)
     77-92-9, Citric acid, uses 7664-38-2, Phosphoric acid, uses
ΤТ
     RL: NUU (Other use, unclassified); USES (Uses)
        (manufacture of re-dispersible blue cobalt zinc silicon oxide pigments)
     155949-59-0, Cobalt silicon zinc oxide
TΤ
     RL: PEP (Physical, engineering or chemical process); PRP (Properties);
     PROC (Process)
        (manufacture of re-dispersible blue cobalt zinc silicon oxide pigments)
ACCESSION NUMBER: 1995:742638 CAPLUS
DOCUMENT NUMBER:
                        123:115526
TITLE:
                        Blue pigment slurries and their manufacture
INVENTOR(S):
                        Mizukami, Tomohito; Funayama, Chihito; Terada, Hiromi;
                        Nishio, Akira
PATENT ASSIGNEE(S):
                         Kasei Optonix, Japan; Dainichiseika Color Chem
SOURCE:
                         Jpn. Kokai Tokkyo Koho, 2 pp.
                         CODEN: JKXXAF
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         Japanese
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
    PATENT NO. KIND DATE APPLICATION NO. DATE

JP 07070466 A 19950314 JP 1993-148931 19930621

RITY APPLN. INFO.: JP 1993-148931 19930621
PRIORITY APPLN. INFO.:
     Title slurries with good re-dispersibility are prepared by wet dispersing
     CoO.ZnO.SiO2 particles in the presence of organic acids or H3PO4.
     Ball-milling 1.2-\mu m CoO .ZnO.SiO2 particles with water and
     citric acid at 110 rpm for 5 h to form 0.4-\mu m particles and setting at
     room temperature for 24 h gave soft and re-dispersible precipitate
L5
    ANSWER 15 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
ΑN
    1992:97969 CAPLUS
```

116:97969

Entered STN: 06 Mar 1992

DN ED

```
TI
     In situ use of gelatin in the preparation of uniform colloidal ferrite
     particles
ΙN
     Siiman, Olavi; Burshteyn, Alexander
PΑ
     Coulter Corp., USA
SO
     U.S., 7 pp.
     CODEN: USXXAM
\mathsf{DT}
     Patent
LA
     English
     ICM B01J013-02
ICS C09D005-23
IC
INCL 252315200
     77-3 (Magnetic Phenomena)
     Section cross-reference(s): 66
FAN.CNT 3
     PATENT NO.
                        KIND DATE APPLICATION NO.
                                                                    DATE
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                        А
                         A 19911105 US 1990-532434 19900604
A1 19911205 CA 1991-2084414 19910531
C 19940104
     US 5062991
PΙ
     CA 2084414
     CA 2084414
                          A1 19911212 WO 1991-US3866
     WO 9119301
                                                                    19910531
         W: AU, CA, JP
         RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, NL, SE
     AU 9181837
                       A 19911231 AU 1991-81837
                                                                    19910531
     AU 654285
                         B2
                               19941103
     EP 532684
                         A1
                               19930324 EP 1991-912637
                  B1
                                                                    19910531
                               19950809
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        R: AT, CH, DE, DK, ES, FR, GB, IT, LI, SE
JP 05509203 T 19931216 JP 1991-513035
ES 2077235 T3 19951116 ES 1991-912637
US 5240640 A 19930831 US 1991-786024
PRAI US 1990-532434 A 19900604
WO 1991-US3866 A 19910531
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19910531
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CLASS
 PATENT NO. CLASS PATENT FAMILY CLASSIFICATION CODES
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                ICM B01J013-02
                 ICS C09D005-23
                 INCL 252315200
                 IPCI B01J0013-02 [ICM, 5]; C09D0005-23 [ICS, 5]
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                         B03C0001-01 [I,A]; H01F0001-032 [I,C*]; H01F0001-06
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                         516/101.000; 252/062.510R; 427/213.350; 428/403.000
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 WO 9119301
                 IPCI
                         B01F0003-00 [ICS,5]; B01J0013-02 [ICS,5]; B32B0005-16
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 AU 9181837
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H01F0001-44 [I,C*]; H01F0001-44 [I,A]
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                          H01F0001-26 [ICM,5]; H01F0001-12 [ICM,5,C*];
                          B01F0003-00 [ICS,5]; B01J0013-02 [ICS,5]; B32B0005-16
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                          H01F0001-44 [I,C*]; H01F0001-44 [I,A]
 JP 05509203
                          H01F0001-26 [ICM,5]; H01F0001-12 [ICM,5,C*];
                   IPCI
                          B01J0013-06 [ICS,5]
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                          G01N0033-543 [I,C*]; G11B0005-133 [I,A]; G11B0005-133 [I,C*]; H01F0001-032 [I,C*]; H01F0001-06 [I,A];
                          H01F0001-11 [I,A]; H01F0001-12 [I,C*]; H01F0001-37
                          [I,A]; HO1FO001-44 [I,A]; HO1FO001-44 [I,C*]
                          516/101.000; 252/062.510R; 252/062.530; 252/062.540;
                  NCL
                          427/213.300; 428/402.000; 428/403.000; 436/526.000;
                          516/100.000
     A 1st solution of KNO3 and KOH or NaNO3 and NaOH, which has been N purged, is
     mixed with a 2nd solution of a Fe2+ salt, divalent metal (Mn2+, Zn2+, Co2+, Ni2+, Ba2+, or mixed Mn2+ and Zn2+) salt, and gelatin, which also has been
     N purged; the mixture of the 2 solns. is swept with N and ripened to form a ferrite hydrosol at a predetd. low temperature for a selected period of time;
     and the hydrosol is washed with the gelatin solution by magnetic separation and
     redispersion to form sep. ferrite particles coated with gelatin.
ST
     colloidal ferrite particle gelatin coated prepn; metal ferrite uniform
     colloidal particle prepn
ΙT
     Gelatins, uses
     RL: PREP (Preparation)
         (in preparation of uniform colloidal metal ferrite particles)
     11129-48-9P, Iron zinc oxide 11138-11-7P, Barium
ΙT
                12023-62-0P, Iron manganese zinc oxide
     (Fe2Mn0.5Zn0.5O4)
                          12052-28-7P, Cobalt ferrite 12063-10-4P, Manganese
                12707-85-6P, Nickel ferrite
     RL: PREP (Preparation)
         (preparation of uniform colloidal particles of, in presence of gelatin)
ACCESSION NUMBER:
                           1992:97969 CAPLUS
DOCUMENT NUMBER:
                           116:97969
TITLE:
                           In situ use of gelatin in the preparation of uniform
                           colloidal ferrite particles
INVENTOR(S):
                           Siiman, Olavi; Burshteyn, Alexander
PATENT ASSIGNEE(S):
                           Coulter Corp., USA
SOURCE:
                           U.S., 7 pp.
                           CODEN: USXXAM
DOCUMENT TYPE:
                           Patent
LANGUAGE:
                           English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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PATENT NO.	KIND DA	TE	APPLICATION NO.		DATE
US 5062991	A 19	911105	US 1990-532434	-	19900604
CA 2084414	A1 19	911205	CA 1991-2084414		19910531
CA 2084414	C 19	940104			
WO 9119301	A1 19	911212	WO 1991-US3866		19910531
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RW: AT, BE, CH,	DE, DK, E	S, FR, GB,	GR, IT, LU, NL, SE		
AU 9181837			AU 1991-81837		19910531
AU 654285	B2 19	941103			
EP 532684	A1 19	930324	EP 1991-912637		19910531
EP 532684	B1 19	950809			
R: AT, CH, DE,	DK, ES, F	R, GB, IT,	LI, SE		
JP 05509203	T 19	931216	JP 1991-513035		19910531
ES 2077235	T3 19	951116	ES 1991-912637		19910531
US 5240640	A 19	930831	US 1991-786024		19911031
PRIORITY APPLN. INFO.:			US 1990-532434	Α	19900604
			WO 1991-US3866	Α	19910531
PRIORITY APPLN. INFO.:					

AB A 1st solution of KNO3 and KOH or NaNO3 and NaOH, which has been N purged, is mixed with a 2nd solution of a Fe2+ salt, divalent metal (Mn2+, Zn2+, Co2+, Ni2+, Ba2+, or mixed Mn2+ and Zn2+) salt, and gelatin, which also has been N purged; the mixture of the 2 solns. is swept with N and ripened to form a ferrite hydrosol at a predetd. low temperature for a selected period of time; and the hydrosol is washed with the gelatin solution by magnetic separation and redispersion to form sep. ferrite particles coated with gelatin.

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L5 ANSWER 16 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
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AN 1991:516761 CAPLUS

DN 115:116761

ED Entered STN: 23 Sep 1991

TI Regeneration of liquid-phase methanol synthesis catalyst

AU Lee, Byung Gwon; Lee, Sunggyu; Kulik, Conrad J.

CS Dep. Chem. Eng., Univ. Akron, Akron, OH, 44325, USA

SO Fuel Science & Technology International (1991), 9(5), 587-612 CODEN: FSCTEG; ISSN: 0884-3759

DT Journal

LA English

CC 45-4 (Industrial Organic Chemicals, Leather, Fats, and Waxes) Section cross-reference(s): 23, 67

OS CASREACT 115:116761

- AB Long-term MeOH production expts. were performed under various reaction environments in order to investigate the relation between catalytic activity and the crystallite size in the MeOH synthesis catalyst. Regeneration expts. were also conducted in order to reduce the crystallite size of the aged catalysts by inducing a metallic-phase redispersion. The drop in the hydrogenation activity was closely linked to the growth in the crystallite size in the catalyst. The crystallite size was reduced successfully by cyclic oxidation-reduction treatments and, as a result, the lost activity in aged catalysts was recovered.
- ST methanol prepn catalyst reactivation; crystallite size hydrogenation catalyst reactivation; oxidn redn catalyst reactivation

IT Hydrogenation catalysts

(cupric oxide-zinc oxide, for carbon oxides to methanol, reactivation of)

IT Crystallites

(reduction in size of, in hydrogenation catalyst reactivation)

IT 1314-13-2, Zinc oxide, uses and miscellaneous

1317-38-0, Cupric oxide, uses and miscellaneous

RL: CAT (Catalyst use); USES (Uses)

(catalysts, for preparation of methanol from synthesis gas, deactivation by crystallite formation in, reactivation by oxidation-reduction treatment for size reduction in)

IT 124-38-9, Carbon dioxide, reactions 630-08-0, Carbon monoxide, reactions

```
RL: RCT (Reactant); RACT (Reactant or reagent)
        (hydrogenation of, methanol from, catalysts for, reactivation by
        crystallite size reduction in)
     67-56-1P, Methanol, preparation
ΙT
     RL: IMF (Industrial manufacture); PREP (Preparation)
        (preparation of, by carbon oxide hydrogenation, catalysts for, reactivation
        by crystallite size reduction in)
ACCESSION NUMBER:
                          1991:516761 CAPLUS
DOCUMENT NUMBER:
                          115:116761
TITLE:
                          Regeneration of liquid-phase methanol synthesis
                          catalyst
AUTHOR(S):
                         Lee, Byung Gwon; Lee, Sunggyu; Kulik, Conrad J.
CORPORATE SOURCE:
                          Dep. Chem. Eng., Univ. Akron, Akron, OH, 44325, USA
SOURCE:
                          Fuel Science & Technology International (1991), 9(5),
                          587-612
                          CODEN: FSCTEG; ISSN: 0884-3759
DOCUMENT TYPE:
                          Journal
LANGUAGE:
                         English
OTHER SOURCE(S):
                          CASREACT 115:116761
     Long-term MeOH production expts. were performed under various reaction
     environments in order to investigate the relation between catalytic
     activity and the crystallite size in the MeOH synthesis catalyst.
     Regeneration expts. were also conducted in order to reduce the crystallite
     size of the aged catalysts by inducing a metallic-phase
     redispersion. The drop in the hydrogenation activity was closely
     linked to the growth in the crystallite size in the catalyst. The
     crystallite size was reduced successfully by cyclic oxidation-reduction
     treatments and, as a result, the lost activity in aged catalysts was
     recovered.
L5
     ANSWER 17 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
ΑN
     1986:115976 CAPLUS
DN
     104:115976
ED
     Entered STN: 05 Apr 1986
TΙ
     Analysis of redispersion properties of suspensions
ΑU
     Gorecki, Mieczyslaw
CS
     Zakl. Chem. Nieorg. Anal., Akad. Med., Poznan, Pol.
SO
     Farmacja Polska (1985), 41(4), 203-4
     CODEN: FAPOA4; ISSN: 0014-8261
DT
     Journal
LΑ
     Polish
CC
     63-5 (Pharmaceuticals)
AB
     Correlations were found between sedimentation coeffs. Vs, thixotropy
     coeffs. \beta, redispersion coeffs. VR, and suspension concns.
     in ZnO or BaSO4 suspensions (particle size 2-9 \mu m, 0.2-0.8
     g/mL) in 1% CM-cellulose [9004-32-4] at 21°. Thus, thixotropy
     coeffs. \beta can be used for the estimation of resuspension properties of
     pharmaceutical suspension after long storage.
ST
     suspension sedimentation thixotropy redispersion
ΙT
     Sedimentation
     Thixotropy
        (of pharmaceutical suspensions)
IT
     Dispersion
        (re-, of pharmaceutical suspension)
IT
     Pharmaceuticals
        (suspensions, redispersion properties of)
ΙT
     1314-13-2, biological studies 7727-43-7
     RL: BIOL (Biological study)
        (suspensions containing, redispersion properties of)
ACCESSION NUMBER:
                         1986:115976 CAPLUS
DOCUMENT NUMBER:
                         104:115976
TITLE:
                         Analysis of redispersion properties of
                         suspensions
AUTHOR(S):
                         Gorecki, Mieczyslaw
CORPORATE SOURCE:
                         Zakl. Chem. Nieorg. Anal., Akad. Med., Poznan, Pol.
```

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SOURCE:
                          Farmacja Polska (1985), 41(4), 203-4
                          CODEN: FAPOA4; ISSN: 0014-8261
DOCUMENT TYPE:
                          Journal
LANGUAGE:
                          Polish
AB
     Correlations were found between sedimentation coeffs. Vs, thixotropy
     coeffs. \beta, redispersion coeffs. VR, and suspension concns.
     in ZnO or BaSO4 suspensions (particle size 2-9 \mu m, 0.2-0.8
     g/mL) in 1% CM-cellulose [9004-32-4] at 21°. Thus, thixotropy coeffs. \beta can be used for the estimation of resuspension properties of
     pharmaceutical suspension after long storage.
L5
     ANSWER 18 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
     1978:465367 CAPLUS
ΑN
DN
     89:65367
ΕD
     Entered STN: 12 May 1984
     Flow equation for coagulated suspensions
ΤI
ΑU
     Cooper, Peter G.; Rayner, John G.; Nicol, Stuart K.
CS
     Cent. Res. Lab., Broken Hill Propr. Co., Ltd., Shortland, Australia
     Journal of the Chemical Society, Faraday Transactions 1: Physical
SO
     Chemistry in Condensed Phases (1978), 74(4), 785-94
     CODEN: JCFTAR; ISSN: 0300-9599
DT
     Journal
LA
     English
CC
     65-1 (General Physical Chemistry)
     An equation for the viscosity at any shear rate was developed from anal.
AΒ
     of the coagulation-redispersion kinetics of a sheared
     suspension. The use of an analogy with chemical reaction kinetics attributes
     some phys. significance to each of the flow consts. in this equation. The
     technique is outlined for the determination of these flow consts. from exptl.
     data, and the equation was fitted to exptl. flow curves for suspensions of
     Fe2O3 and ZnO in H2O.
ST
     flow shear coagulation suspension; ferric oxide suspension flow;
     zinc oxide suspension flow
ΙT
     Coagulation
         (-redispersion, of sheared suspensions, flow in relation to)
ΙT
     Flow
        (equations of, for sheared coagulated suspensions)
ΙT
     Suspensions
        (flow of sheared coagulated, equations for)
ΙT
     1309-37-1, properties 1314-13-2, properties
     RL: PRP (Properties)
        (flow of coagulated suspensions of, theor. flow equation in relation
        to)
ACCESSION NUMBER:
                          1978:465367 CAPLUS
DOCUMENT NUMBER:
                          89:65367
TITLE:
                          Flow equation for coagulated suspensions
AUTHOR(S):
                          Cooper, Peter G.; Rayner, John G.; Nicol, Stuart K.
CORPORATE SOURCE:
                          Cent. Res. Lab., Broken Hill Propr. Co., Ltd.,
                          Shortland, Australia
SOURCE:
                          Journal of the Chemical Society, Faraday Transactions
                          1: Physical Chemistry in Condensed Phases (1978),
                          74(4), 785-94
CODEN: JCFTAR; ISSN: 0300-9599
DOCUMENT TYPE:
                          Journal
LANGUAGE:
                          English
     An equation for the viscosity at any shear rate was developed from anal.
     of the coagulation-redispersion kinetics of a sheared
     suspension. The use of an analogy with chemical reaction kinetics attributes
     some phys. significance to each of the flow consts. in this equation. The
     technique is outlined for the determination of these flow consts. from exptl.
     data, and the equation was fitted to exptl. flow curves for suspensions of
     Fe2O3 and ZnO in H2O.
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ANSWER 19 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

1970:124991 CAPLUS

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DN
     72:124991
ED
     Entered STN: 12 May 1984
     Behavior of stabilized lotions under thermal conditions
TΙ
ΑU
     Rohdewald, Peter
     Inst. Pharm. Chem., Westfael. Wilhelm-Univ., Muenster, Fed. Rep. Ger.
CS
SO
     Pharmazie (1969), 24(12), 773-5
     CODEN: PHARAT; ISSN: 0031-7144
DT
     Journal
LA
     German
CC
     62 (Essential Oils and Cosmetics)
AB
     Zinc oxide-talc suspensions stabilized with
     CM-cellulose, methylcellulose, propylene glycol alginate, bentonite, or
     self-emulsifying wax were not stable on storage at 50° for 40 days.
     The organic stabilizers caused caking of the sediment so that it could not be
     redispersed. The wax-stabilized suspension was particularly unstable.
     Heating at 95° for 2 hr did not decrease the stability of the
     suspensions and improved the redispersion of the sediment.
     Brookfield viscosity detns. gave an accurate measure of the condition of
     the suspensions.
     zinc oxide suspension stabilized; talc suspension
ST
     stabilized; suspension stabilized talc ZnO; stabilized
     suspension talc ZnO
     Lotions
IT
        (sediment caking in, stabilizer effect on)
IT
     Bentonite, uses and miscellaneous
     Waxes
     RL: BIOL (Biological study)
        (stabilizers, sediment caking in lotions in relation to)
     1,2-Propanediol, alginate
ΙT
     RL: BIOL (Biological study)
        (stabilizers, sediment caking in lotions in relation to)
                 9004-67-5 9005-37-2, Alginic acid, ester with
ΙT
     9000-11-7
     1,2-propanediol
     RL: BIOL (Biological study)
        (stabilizers, sediment caking in lotions in relation to)
ACCESSION NUMBER:
                         1970:124991 CAPLUS
DOCUMENT NUMBER:
                         72:124991
TITLE:
                         Behavior of stabilized lotions under thermal
                         conditions
AUTHOR(S):
                         Rohdewald, Peter
CORPORATE SOURCE:
                         Inst. Pharm. Chem., Westfael. Wilhelm-Univ., Muenster,
                         Fed. Rep. Ger.
SOURCE:
                         Pharmazie (1969), 24(12), 773-5
                         CODEN: PHARAT; ISSN: 0031-7144
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         German
     Zinc oxide-talc suspensions stabilized with
     CM-cellulose, methylcellulose, propylene glycol alginate, bentonite, or
     self-emulsifying wax were not stable on storage at 50° for 40 days.
     The organic stabilizers caused caking of the sediment so that it could not be
     redispersed. The wax-stabilized suspension was particularly unstable.
     Heating at 95° for 2 hr did not decrease the stability of the
     suspensions and improved the redispersion of the sediment.
     Brookfield viscosity detns. gave an accurate measure of the condition of
     the suspensions.
L5
     ANSWER 20 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
     1969:48879 CAPLUS
ΑN
DN
     70:48879
ED
     Entered STN: 12 May 1984
ΤI
     Continuous hydrolysis of animal fats and oils
ΑU
     Jenson, Victor G.; Jeffreys, Godfrey V.; Edwards, Reginald Eric
CS
     Univ. Birmingham, Birmingham, UK
     Industrie Chimique Belge (1967), 32(Spec. No.), 694-9
SO
     CODEN: ICBEAJ; ISSN: 0019-9052
```

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DT
     Journal
LA
     English
CC
     45 (Fats and Waxes)
     A 3-in.-diameter, 8-ft.-long hydrolyzer column was set up and 16 expts. were
     done at various flow rates with beef tallow with or without ZnO
     as a catalyst. Four expts. were also done with grid plates in the column.
     The reaction mechanism corresponded to a 2nd-order description. Equations
     were set up and the exptl. data integrated. By addition of 0.3% _{200} , a 70% increase in conversion was obtained. The exptl. runs with grid
     plates could not be simulated on the computer due to the effect of
     coalescence and redispersion on the plates giving unknown
     mass-transfer coeffs. and interfacial areas, but the plates improved the
     column performance.
     fat hydrolysis continuous; oil hydrolysis continuous; animal fat
    hydrolysis; beef tallow hydrolysis; hydrolysis fat oil
     Kinetics of saponification
ΙΤ
         (of tallow)
ΙT
     Saponification
        (of tallow in long columns)
TΤ
     Tallow
     RL: USES (Uses)
         (saponification of beef, in long columns)
IT
     Saponification catalysts
        (zinc oxide, for tallow)
IT
     1314-13-2, uses and miscellaneous
     RL: CAT (Catalyst use); USES (Uses)
         (catalysts, for saponification of tallow)
ACCESSION NUMBER:
                          1969:48879 CAPLUS
DOCUMENT NUMBER:
                          70:48879
TITLE:
                          Continuous hydrolysis of animal fats and oils
AUTHOR(S):
                          Jenson, Victor G.; Jeffreys, Godfrey V.; Edwards,
                          Reginald Eric
CORPORATE SOURCE:
                          Univ. Birmingham, Birmingham, UK
SOURCE:
                          Industrie Chimique Belge (1967), 32(Spec. No.), 694-9
                          CODEN: ICBEAJ; ISSN: 0019-9052
DOCUMENT TYPE:
                          Journal
LANGUAGE:
                          English
     A 3-in.-diameter, 8-ft.-long hydrolyzer column was set up and 16 expts. were
     done at various flow rates with beef tallow with or without ZnO
     as a catalyst. Four expts. were also done with grid plates in the column.
     The reaction mechanism corresponded to a 2nd-order description. Equations
     were set up and the exptl. data integrated. By addition of 0.3\% ZnO , a 70% increase in conversion was obtained. The exptl. runs with grid
     plates could not be simulated on the computer due to the effect of
     coalescence and redispersion on the plates giving unknown
     mass-transfer coeffs. and interfacial areas, but the plates improved the
     column performance.
     ANSWER 21 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN
L5
AN
     1967:509296 CAPLUS
DN
     67:109296
OREF 67:20643a,20646a
ED
     Entered STN: 12 May 1984
     Poly(vinyl alcohol) adhesives
TI
PΑ
     Kurashiki Rayon Co., Ltd.
     Brit., 3 pp. CODEN: BRXXAA
SO
DT
     Patent
LA
     English
IC
     C08F
CC
     36 (Plastics Manufacture and Processing)
FAN.CNT 1
     PATENT NO.
                         KIND
                                             APPLICATION NO.
                                 DATE
                                 -----
                          ____
                                              -----
                                 19670831 GB 1964-32735
PT
     GB 1081413
                                                                      19640811
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DE 1594071
                                             DE
     US 3365408
                                 19680123
                                             US 1964-387203 19640803
PRAI JP
                                 19630812
CLASS
 PATENT NO.
                CLASS PATENT FAMILY CLASSIFICATION CODES
 GB 1081413
                        C08F
                 IPCI
                        COSE
                        C08L0033-00 [N,C*]; C08L0033-02 [N,A]; C09J0129-00
                 IPCR
                        [I,C*]; C09J0129-04 [I,A]
 US 3365408
                        C08L0033-00 [N,C*]; C08L0033-02 [N,A]; C09J0129-00
                 IPCR
                         [I,C*]; C09J0129-04 [I,A]
                 NCL
                        524/401.000; 524/430.000; 524/431.000; 524/432.000;
                        524/433.000; 524/434.000; 524/435.000; 524/436.000;
                        524/437.000; 524/446.000; 524/447.000; 524/503.000;
                        525/056.000; 525/061.000
     The title compns. are prepared from a polyvalent metal oxide or hydroxide
     and a poly(vinyl alc.) (I) derivative containing 5-70 mole% CO2H groups or a
mixture
     of I and a polymer containing 5-70 mole% CO2H groups. Thus, an aqueous
solution
     containing 5% I and 5% poly(acrylic acid) was heated at 90% for 1 hr., 2.0%
     weight ZnO added and thoroughly dispersed, and the mixture allowed to
     stand for 3 hrs. After redispersion, the composition was spread on a
     poly(vinyl chloride) plate and heat-treated 3 min. at 120°, giving
     a film with water solubility 34.6%. An untreated film had a solubility of
     pure I, 100%. This adhesive gave an initial adhesion of 1.0 kg. between a
     corrugated liner and center board after 2 sec. at 120°. CaO and
     clay were also used, and oxides and hydroxides of Mg, Zn, Cu, Al, and Fe
     are claimed. Saponified vinyl acetate (II)-acrylate, II-methacrylate,
     II-acrylamide copolymer, or II-acrylonitrile copolymers, and I etherified
     with glycolic acid or acetalized with glyoxylic acid are also claimed.
     Saponified polyacrylamide or polyacrylonitrile can also be used as a mixture
     with II. These adhesives have improved water resistance, and are used for
     manufacturing corrugated paperboard boxes. POLYVINYL ALC ADHESIVE; POLYACRYLIC ACID; ADHESIVE POLYVINYL ALC;
ST
     PAPERBOARD ADHESIVE
ΙT
     Paperboard
        (adhesives for corrugated, vinyl alc. polymer solns. containing carboxylic
        acid polymers and metal oxides as)
IT
     Paper
        (as catalyst support for oxygen removal from, with clay)
ΙT
     Clays
     RL: USES (Uses)
        (coating with, of paper, adhesives for)
IT
     Adhesives, preparation
        (from vinyl alc. polymer solns. containing carboxylic acid polymers and
        metal oxides)
ΙT
     Coating process
        (of paper with clay, vinyl alc. polymers adhesives for)
     9002-89-5, uses and miscellaneous
IT
     RL: USES (Uses)
        (adhesives from carboxylic acid polymers, metal oxides and)
ΙT
     1305-78-8, uses and miscellaneous 1314-13-2, uses and miscellaneous
     RL: USES (Uses)
        (adhesives from vinyl alc. polymer solns. containing carboxylic acid
        polymers and)
TT
     9003-01-4
     RL: USES (Uses)
        (adhesives from vinyl alc. polymer solns. containing metals oxides and)
ACCESSION NUMBER:
                         1967:509296 CAPLUS
DOCUMENT NUMBER:
                         67:109296
ORIGINAL REFERENCE NO.: 67:20643a,20646a
TITLE:
                         Poly(vinyl alcohol) adhesives
```

PATENT ASSIGNEE(S): Kurashiki Rayon Co., Ltd.

SOURCE: Brit., 3 pp.
CODEN: BRXXAA

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
GB 1081413		19670831	GB 1964-32735	19640811
DE 1594071 US 3365408		19680123	DE US 1964-387203	19640803

PRIORITY APPLN. INFO.: JP 19630812

AB The title compns. are prepared from a polyvalent metal oxide or hydroxide

and a poly(vinyl alc.) (I) derivative containing 5-70 mole% CO2H groups or a mixture

of I and a polymer containing 5-70 mole% ${\tt CO2H}$ groups. Thus, an aqueous solution

containing 5% I and 5% poly(acrylic acid) was heated at 90% for 1 hr., 2.0% weight ZnO added and thoroughly dispersed, and the mixture allowed to stand for 3 hrs. After redispersion, the composition was spread on a poly(vinyl chloride) plate and heat-treated 3 min. at 120°, giving a film with water solubility 34.6%. An untreated film had a solubility of 90%, and

pure I, 100%. This adhesive gave an initial adhesion of 1.0 kg. between a corrugated liner and center board after 2 sec. at 120°. CaO and clay were also used, and oxides and hydroxides of Mg, Zn, Cu, Al, and Fe are claimed. Saponified vinyl acetate (II)-acrylate, II-methacrylate, II-acrylamide copolymer, or II-acrylonitrile copolymers, and I etherified with glycolic acid or acetalized with glyoxylic acid are also claimed. Saponified polyacrylamide or polyacrylonitrile can also be used as a mixture with II. These adhesives have improved water resistance, and are used for manufacturing corrugated paperboard boxes.

L5 ANSWER 22 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

AN 1957:88183 CAPLUS

DN 51:88183 OREF 51:15967b-d

ED Entered STN: 22 Apr 2001

TI Applications of dry vinyl ester dispersions in coating technology

AU Grunau, E. B.

SO Farbe u. Lack (1957), 63, 345-6

DT Journal

LA Unavailable

CC 26 (Paints, Varnishes, Lacquers, and Inks)

Vinyl acetate powders are either difficult to redisperse or do not form a AB continuous film after mixing with water. An improved dry dispersion can be prepared by coating the spray dried powder at once with a fine particle-size pigment. Poly(vinyl acetate) latexes containing 12-15% plasticizer (based on total solids) and vinyl copolymers, having a particle size of 0.7 to 1.6 μ , give the best results. Fine zinc oxides are especially suitable pigments and are mixed into the dispersion by tangential blowing. From 25 to 30 parts of pigment per 100 parts emulsion solids are necessary to prevent caking. Addnl. pigments and fillers can later be blended with the dry powder. From 2 to 3.5% of solvents, glycols and ketones, should be added to the water used for redispersion of the powder to improve film For viscosity control of the dispersion, dry poly(vinyl alc.) can be blended with the pigments. Dry dispersions are economical to manufacture and the paint films are equal in properties to those produced from conventional latex paints.

IT Coating(s)

(from vinyl ester dispersion in dry state) ACCESSION NUMBER: 1957:88183 CAPLUS

DOCUMENT NUMBER: 51:88183
ORIGINAL REFERENCE NO.: 51:15967b-d

TITLE: Applications of dry vinyl ester dispersions in coating

technology Grunau, E. B.

SOURCE: Farbe u. Lack (1957), 63, 345-6

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

Vinyl acetate powders are either difficult to redisperse or do not form a continuous film after mixing with water. An improved dry dispersion can be prepared by coating the spray dried powder at once with a fine particle-size pigment. Poly(vinyl acetate) latexes containing 12-15% plasticizer (based on total solids) and vinyl copolymers, having a particle size of 0.7 to 1.6 $\mu\text{,}$ give the best results. Fine zinc oxides are especially suitable pigments and are mixed into the dispersion by tangential blowing. From 25 to 30 parts of pigment per 100 parts emulsion solids are necessary to prevent caking. Addnl. pigments and fillers can later be blended with the dry powder. From 2 to 3.5% of solvents, glycols and ketones, should be added to the water used for redispersion of the powder to improve film formation. For viscosity control of the dispersion, dry poly(vinyl alc.) can be blended with the pigments. Dry dispersions are economical to manufacture and the paint films are equal in properties to those produced from conventional latex paints.

L5 ANSWER 23 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN

AN 1950:26284 CAPLUS

DN 44:26284

AUTHOR(S):

OREF 44:5138i,5139a-e

ED Entered STN: 22 Apr 2001

TI Structure of vulcanized latex

AU Humphreys, N. C. H.; Wake, W. C.

CS Research Assoc. Brit. Rubber Manufrs., Croydon, UK

SO Transactions, Institution of the Rubber Industry (1950), 25, 334-45

CODEN: TIRIA2; ISSN: 0371-7968

DT Journal

LA Unavailable

CC 30 (Rubber and Other Elastomers)

AB The hydrocarbon phase of latex can be regarded as an oil phase, and withdrawal of most of the aqueous phase by evaporation leaves an unstable condition

until the orientation of the interracial material changes. This takes place by a process analogous to phase inversion, whereby the remaining water diffuses through the hydrocarbon. If the protein is replaced by saponin, this reorientation is evident. Thus, when the water is still at least 1% during drying, the film can be redispersed by soaking in water; below 1% redispersion becomes impossible, and at this point the saponin mols. orient themselves to form a water in-oil type of structure. This phase inversion is evident also by a sudden increase in elec. resistivity when the water content decreases to a sufficiently low value. Latex was recreamed 4 times in the presence of Na laurate to displace the protein, then vulcanized, strained, and dried at room temperature on glass plates. In another experiment, the soap was extracted by EtOH and drying was

then

completed. All films had the same strength as that of corresponding normally vulcanized films. Hence cohesion must be by ordinary chemical forces, i.e., either by primary-valence bonds or by secondary bonds or van der Waals' forces between rubber mols. Since inhibition of a solvent destroys only secondary valence bonds, the 2 possibilities were tested by swelling dried vulcanized films in C6H6. Swelling maxima were obtained which indicated cross-linking, but the phys. properties indicated the absence of primary-valence bonds between particles and their presence between particles. Hence the cohesion is attributable to van der Waals' forces between the outer mols. of the particles. All these expts. and considerations disprove theoretically and experimentally the view of

Hauser (Latex, p. 130) that adsorbed protein and other nonrubber components bond the rubber particles. Extraction with CHCl3 of acetone-extracted

vulcanized film gave only 0.3% extract of rubber of low mol. weight; hence, the rubber within the latex particles is vulcanized, i.e., practically completely cross-linked. To a latex mixture prepared from 167 cc. 60% latex., 1 g. ZnO, 1 g. Na diethyldithiocarbamate, 0.25 g. casein, and $0.5~\mathrm{g}$. NaOH, were added 2, 2.5, 3, 4, 10, and $50~\mathrm{g}$., resp., of S, and the mixts. were vulcanized and centrifuged. In no case was the combined S over 1.8%, based on the rubber. The stress-strain curves of the dried vulcanized films show very low moduli for low extensions after which they increase rapidly to the breaking point. The structure of films from vulcanized latex which is proposed is discussed in its relation to the phys. properties of the films and in the light of modern theories of elasticity.

IT Films

(from rubber, phys. properties and structure of)

ΙT Cohesion

(of rubber mols.)

TΤ Molecules

(orientation and structure of, of rubber)

IT Rubber

(structure of vulcanized latex)

ACCESSION NUMBER: 1950:26284 CAPLUS

DOCUMENT NUMBER: 44:26284

ORIGINAL REFERENCE NO.: 44:5138i,5139a-e

Structure of vulcanized latex TITLE: Humphreys, N. C. H.; Wake, W. C. AUTHOR(S):

CORPORATE SOURCE: Research Assoc. Brit. Rubber Manufrs., Croydon, UK SOURCE: Transactions, Institution of the Rubber Industry

(1950), 25, 334-45 CODEN: TIRIA2; ISSN: 0371-7968

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

The hydrocarbon phase of latex can be regarded as an oil phase, and withdrawal of most of the aqueous phase by evaporation leaves an unstable

until the orientation of the interracial material changes. This takes place by a process analogous to phase inversion, whereby the remaining water diffuses through the hydrocarbon. If the protein is replaced by saponin, this reorientation is evident. Thus, when the water is still at least 1% during drying, the film can be redispersed by soaking in water; below 1% redispersion becomes impossible, and at this point the saponin mols. orient themselves to form a water in-oil type of structure. This phase inversion is evident also by a sudden increase in elec. resistivity when the water content decreases to a sufficiently low value. Latex was recreamed 4 times in the presence of Na laurate to displace the protein, then vulcanized, strained, and dried at room temperature on glass plates. In another experiment, the soap was extracted by EtOH and drying was

then

acetone-extracted

completed. All films had the same strength as that of corresponding normally vulcanized films. Hence cohesion must be by ordinary chemical forces, i.e., either by primary-valence bonds or by secondary bonds or van der Waals' forces between rubber mols. Since inhibition of a solvent destroys only secondary valence bonds, the 2 possibilities were tested by swelling dried vulcanized films in C6H6. Swelling maxima were obtained which indicated cross-linking, but the phys. properties indicated the absence of primary-valence bonds between particles and their presence between particles. Hence the cohesion is attributable to van der Waals' forces between the outer mols. of the particles. All these expts. and considerations disprove theoretically and experimentally the view of Hauser (Latex, p. 130) that adsorbed protein and other nonrubber components bond the rubber particles. Extraction with CHC13 of

vulcanized film gave only 0.3% extract of rubber of low mol. weight; hence, the

rubber within the latex particles is vulcanized, i.e., practically completely cross-linked. To a latex mixture prepared from 167 cc. 60% latex., 1 g. ZnO, 1 g. Na diethyldithiocarbamate, 0.25 g. casein, and 0.5 g. NaOH, were added 2, 2.5, 3, 4, 10, and 50 g., resp., of S, and the mixts. were vulcanized and centrifuged. In no case was the combined S over 1.8%, based on the rubber. The stress-strain curves of the dried vulcanized films show very low moduli for low extensions after which they increase rapidly to the breaking point. The structure of films from vulcanized latex which is proposed is discussed in its relation to the phys. properties of the films and in the light of modern theories of elasticity.

L5 ANSWER 24 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN ΑN 1947:12716 CAPLUS DN 41:12716 OREF 41:2587b-f Entered STN: 22 Apr 2001 ED ΤI The settling of paints ΑU Rabate, H. SO Peintures, Pigments, Vernis (1946), 22, 213-18,253-8 CODEN: PPVEAF; ISSN: 0031-4102 DT Journal Unavailable LA CC 26 (Paints, Varnishes, and Lacquers) AB The factors influencing the settling of a paint are: the d. of the solid and liquid phases; the size, distribution, and shape of the pigment or filler particles; the nature of the solid surface (compactness, porosity, chemical reactivity, moisture adsorption, hydrophilic or organophilic character); the viscosity and nature (chemical stability, colloidal character, acidity, polarity, wetting, and dispersing power) of the liquid phase; the weight ratio solid/ liquid; the technique employed in grinding and mixing; the conditions of storage. The sedimentation rule of Stokes has only limited application; the relation of viscosity to the pigment volume of a paint is expressed by the rule of Wolff and Zeidler (C.A. 23, 7878). The settling rate is influenced by the thickness of the adsorbed film-around the pigment particles. Well-wetted pigments cake rapidly and hard, while poorly wetted ones form soft, loose deposits. Electrostatic charges of the suspended solids have little influence on settling; rigidity of a sediment can favorably influence the settling rate (cf. McMillen and Glaser, C.A. 32, 8167.2). Generally, the sediments in colloidal vehicles are harder than those in non-colloidal ones, hardness increasing with vehicle viscosity. Ease of redispersion of a sediment depends upon how thoroughly the solids are wetted by the vehicle. Heat and certain wetting agents or stabilizers influence ease of redispersion and suspension stability. For more rapid evaluation, the paint may be thinned, or centrifuged hot, or subjected to heat and centrifugation alternately. The sediment consistency can be measured by a needle penetrometer or the ease of torsion of an immersed plunger. The settling of paints can be reduced by selecting fillers of low d., small particle size, and particles of suitable shape, by using a vehicle of high viscosity, or thickened with metallic soaps, by selecting media capable of gelation by adding reactive pigments (e.g. ZnO), and by the addition of dispersing or wetting agents. 26 references. ΙT Particles (concentration, shape and size of, of paint fillers, effect on settling of paint) ΙT Wetting agents (effect on redispersion of sediments in settled paints) ΙT Grinding (effect on settling of paints) ΙΤ Consistency (measurement of, of paint sediment) IT Sedimentation (of paints)

IΤ

Electric charge

(on pigments and paint fillers, effect on settling of paints) ΙT Dispersion (re-, of sediments in paints) TT Paint (sedimentation in) ΙT Pigments (settling of paints in relation to) ACCESSION NUMBER: 1947:12716 CAPLUS DOCUMENT NUMBER: 41:12716 ORIGINAL REFERENCE NO.: 41:2587b-f TITLE: The settling of paints AUTHOR(S): Rabate, H. SOURCE: Peintures, Pigments, Vernis (1946), 22, 213-18,253-8 CODEN: PPVEAF; ISSN: 0031-4102 DOCUMENT TYPE: LANGUAGE: Unavailable The factors influencing the settling of a paint are: the d. of the solid and liquid phases; the size, distribution, and shape of the pigment or filler particles; the nature of the solid surface (compactness, porosity, chemical reactivity, moisture adsorption, hydrophilic or organophilic character); the viscosity and nature (chemical stability, colloidal character, acidity, polarity, wetting, and dispersing power) of the liquid phase; the weight ratio solid/ liquid; the technique employed in grinding and mixing; the conditions of storage. The sedimentation rule of Stokes has only limited application; the relation of viscosity to the pigment volume of a paint is expressed by the rule of Wolff and Zeidler (C.A. 23, 7878). The settling rate is influenced by the thickness of the adsorbed film-around the pigment particles. Well-wetted pigments cake rapidly and hard, while poorly wetted ones form soft, loose deposits. Electrostatic charges of the suspended solids have little influence on settling; rigidity of a sediment can favorably influence the settling rate (cf. McMillen and Glaser, C.A. 32, 8167.2). Generally, the sediments in colloidal vehicles are harder than those in non-colloidal ones, hardness increasing with vehicle viscosity. Ease of redispersion of a sediment depends upon how thoroughly the solids are wetted by the vehicle. Heat and certain wetting agents or stabilizers influence ease of redispersion and suspension stability. For more rapid evaluation, the paint may be thinned, or centrifuged hot, or subjected to heat and centrifugation alternately. The sediment consistency can be measured by a needle penetrometer or the ease of torsion of an immersed plunger. The settling of paints can be reduced by selecting fillers of low d., small particle size, and particles of suitable shape, by using a vehicle of high viscosity, or thickened with metallic soaps, by selecting media capable of gelation by adding reactive pigments (e.g. ZnO), and by the addition of dispersing or wetting agents. 26 references. ANSWER 25 OF 26 CAPLUS COPYRIGHT 2007 ACS on STN L51927:11404 CAPLUS AN DN 21:11404 OREF 21:1375i,1376a-e Entered STN: 16 Dec 2001 ED ΤI Effects of some organic acids on the rate of the vulcanization of rubber ΑU Mitchell, P. H. SO Caoutchouc et la Gutta-Percha (1927), 24, 13,440-3 CODEN: CAGUAC; ISSN: 0366-5224 DT Journal LA Unavailable 30 (Rubber and Allied Substances) CC A survey of the literature on the effect of acids and resins on the rate of vulcanization of rubber shows a general lack of information regarding the relative effects of individual acids on the rate of vulcanization and on the phys. properties of the vulcanizates. A study was therefore made. of the effects of some of the acids in the resin of Hevea rubber and certain other acids with m. ps. not above the temperature of vulcanization. To rubber-S (92.5-7.5) mixts., the rubber of which had previously been extracted

successively with H2O and with Me2CO, were added the individual acids in amts. equivalent in the H2O-soluble acids to the total-acidity of the rubber,

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in the H2O-insol. acids to the acidity of the resin. Parallel series were in turn prepared with and without just enough ZnO (0.1767 g.) to neutralize the acid. The effects on the rate of vulcanization were judged by comparing the times to reach the maximum tensile strength, which was called the optimum cure. The long time of cure and the small quantities of acids made it difficult in some cases to ascribe the small changes in the time of optimum cure to the acids alone, and 15 min. differences in cure are of little significance. The effects of the acids are most readily shown by the following data, which give the % acid added to the extracted rubber, the time of cure to reach the maximum tensile strength of the mixts. without ZnO and the time for the corresponding mixts. containing ZnO, resp.: unextd. rubber (no acid), -, 225, -; extracted rubber (no acid), , 180, 210; Me2CO extract, 3.005, 195, ; stearic acid, 1.18, 180, 210; oleic acid, 1.177, 180, 195; linoleic acid, 1.168, 180, 180; palmitic acid, 1.06, 225, 225; lactic acid, 0.40, 225, 255; malic acid, 0.298, 225, 225; levulinic acid, 0.515, 240, 240; butyric acid, 0.392, 195, 210; caproic acid, 0.478, 225, 240; isocaproic acid, 0.478, 210, 180; caprylic acid, 0.641, 210, 150; benzoic acid, 0.504, 225, ; salicylic acid, 0.569, 300, -. The retardant action of the rubber resins is not in accord with the earlier work of Whitby and of Stevens, but is confirmed by the retardant action observed on adding the individual resin acids to the extracted rubber mixture Since the extracted rubber mixture to which the

Me2CO extract was returned vulcanized more rapidly than the unextd. rubber mixture and was of poorer quality, there must be a H2O-soluble acid with a retardant action, or a loss of volatile Me2CO-soluble acid, or some phys. cause such as incomplete redispersion of the resin. The increase in the time of cure of the extracted rubber mixture on addition of ZnO is not easily explained, but may be due to the effect of the ZnO on the phys. properties, since the mixture containing ZnO was much the superior. The notable retarding effect of lactic acid on the rate of vulcanization and the fact that it is a good coagulant favors the theory of Whitby and Cambron (C. A. 17, 3808). In further expts. oleic, butyric and caproic acids were added to an extracted rubber-S-ZnO (92.5-7.5-10) mixture, in which case butyric acid had no effect (time of cure 195 min. with and without acid), whereas oleic acid and caproic acid accelerated the rate of vulcanization (time of cure 150 min. each). The results indicate that when there is merely enough ZnO to neutralize the acid, the soaps have no noticeable accelerating action, and that with excess ZnO the chief function of the resin is to improve the dispersion of the pigment. Twenty-six references are included.

ITAcids

(effect of organic, on vulcanization of rubber)

Rubber

(vulcanization of, effect of organic acids on)

ACCESSION NUMBER: 1927:11404 CAPLUS

DOCUMENT NUMBER: 21:11404

ORIGINAL REFERENCE NO.: 21:1375i,1376a-e

TITLE: Effects of some organic acids on the rate of the

vulcanization of rubber

AUTHOR(S): Mitchell, P. H.

SOURCE: Caoutchouc et la Gutta-Percha (1927), 24, 13,440-3

CODEN: CAGUAC; ISSN: 0366-5224

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

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ΑN
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     Entered STN: 16 Dec 2001
ΤI
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     Mitchell, P. H.
     Caoutchouc et la Gutta-Percha (1926), 23, 13,340-2
SO
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